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(54) Title: METHODS OF INHIBITING OR ENHANCING THE  $TGF\beta$ -SMAD SIGNALING PATHWAY

#### (57) Abstract

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Signal transduction by the TGF- $\beta$  family involves sets of receptor serine/threonine kinases, Smad proteins that act as receptor substrates, and Smad-associated transcription factors that target specific genes. Discrete structural elements were identified that dictate the selective interactions between receptors and Smads and between Smads and transcription factors in the TGF- $\beta$  and BMP pathways. A cluster of four residues in the L45 loop of the type I receptor kinase domain, and a matching set of two residues in the L3 loop of the Smad C-terminal domain establish the specificity of receptor-Smad interactions. A cluster of residues in the highly exposed  $\alpha$ -helix 2 of the Smad C-terminal domain specify the interaction with the DNA-binding factor Fast1 and, as a result, the gene responses mediated by the pathway. By establishing specific interactions, these determinants keep the TGF- $\beta$  and BMP pathways segregated from each other.

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# METHODS OF INHIBITING OR ENHANCING THE $TGF\beta$ -SMAD SIGNALING PATHWAY

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#### BACKGROUND OF THE INVENTION

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#### Cross-Referenced to Related Applications

This non-provisional application claims the benefit of priority of provisional applications United States Serial Numbers 60/052,774 filed on July 1, 1997 and 60/065,113 filed on November 12, 1997.

#### Federal Funding Legend

This invention was created in part using funds from the National Institutes of Health under grant R37-CA34610. The federal government, therefore, has certain rights in this invention.

#### Field of the Invention

The present invention relates generally to the fields of molecular biology and cellular biology of cytokines. More specifically, the present invention relates to a methods of inhibiting or enhancing the TGF-β-Smad signaling pathway.

#### Description of the Related Art

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The TGF-β family of polypeptide growth factors regulate cell division, differentiation, motility, adhesion and death in virtually all metazoan tissues<sup>39,44,46,51,53,56</sup>. Members of this family include the TGF-\u03b3s, the activins, the bone morphogenetic proteins (BMPs) and other related factors. Signal transduction by these factors involves three classes of molecules: a family of membrane receptor serine/threonine kinases, a family cytoplasmic proteins, the Smad family, that serve as substrates for these receptors, and nuclear DNA-binding factors that associate with Smads forming transcriptional complexes<sup>43,52</sup>. Signaling is initiated by binding of the growth factor to a specific pair of receptor kinases, an event that induces the phosphorylation and activation of one kinase, known as the "type I receptor", by the other kinase or "type II receptor"65. The activated type I receptor phosphorylates a subset of Smads, known as "receptor-regulated Smads" (R-Smads), which then move into the nucleus<sup>43.52</sup>. On their way to the nucleus, R-Smads associate with the related protein Smad49, a tumor suppressor gene product. In the nucleus, this complex may associate with specific DNA-binding proteins that direct it to the regulatory region of target genes. The first identified Smad-associated DNA-binding factor was the forkhead family member Fast1, which mediates activation of Mix.2 in response to activin-type signals during Xenopus embryogenesis<sup>36</sup>. The integrity of this signaling network is essential for normal development and tissue homeostasis, and its disruption by

mutation underlies several human inherited disorders and cancer<sup>43,52</sup>.

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Because of the diversity of processes controlled by different TGF- $\beta$  family members, there is an intense interest in elucidating the basis for the specificity of their signal transduction pathways. The TGF-β and activin type I receptors, which have identical domains<sup>31,60</sup>, interact kinase nearly with and phosphorylate Smad2 (or the closely related Smad3)16,40,30,54,8 which then interacts with DNA-binding factors such as Fast 134,33,49 The BMP receptors interact with Smadl (or the closely related Smads 5, 8 or, in Drosophila, Mad) 35.40.11.14.18.10 which do not recognize Fast1<sup>36</sup>. Although the TGF-β and BMP pathways are well segregated from each other, their receptors and R-Smads are structurally very similar. The specificity of the receptor and Smad interactions in each pathway may therefore be dictated discrete structural elements.

The Smad4/DPC4 tumor suppressor¹ is inactivated in nearly one half of pancreatic carcinomas² and to a lesser extent in a other cancers²-⁴. Smad4/DPC4, and the related tumor suppressor Smad2, belong to the Smad family of proteins which mediate TGFβ/activin/bone morphogenetic protein (BMP)-2/4 cytokine superfamily signaling from the receptor serine/threonine protein kinases at the cell surface to the nucleus⁵-⁻. Smad proteins, which get phosphorylated by the activated receptor, propagate the signal, in part, through homo-oligomeric and hetero-oligomeric interactions²-¹³. Smad4/DPC4 plays a central role as it is the shared hetero-oligomerization partner of the other Smads. The conserved C-terminal domains of Smads are sufficient for inducing

most of the ligand-specific effects, and are the primary targets of tumorigenic inactivation.

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The conserved C-terminal domain of Smad family members is the likely effector domain, whereas the conserved Nterminal domain is the likely negative regulator of activity<sup>14</sup>. in a Smad4/DPC4-/overexpressed cell line. Smad4/DPC4 C-terminal domain activates the transcription TGF-\(\beta\) responsive genes and results in growth arrest in a ligandindependent manner, paralleling the effects of the TGF-β ligand9. In addition, microinjection of mRNAs encoding the C-terminal domain of Smad2 into Xenopus embryos can induce a mesoderm response that mimics the effects of the full-length protein 16. Furthermore, the Smad4/DPC4-C-terminal domain fused to a heterologous DNA-binding domain can activate gene expression from a reporter construct<sup>14</sup>. Consistent with the Smad C-terminal domain being the main effector domain, the majority (10 out of 13) of the tumorigenic missense mutations in Smad4/DPC4 and Smad2, as well as mutations isolated from Drosophila and C elegans genetic screens map to the C-terminal domain.

The prior art is deficient in the lack of effective means of inhibiting or enhancing the  $TGF-\beta$ -Smad signaling pathway. The present invention fulfills this longstanding need and desire in the art.

#### **SUMMARY OF THE INVENTION**

It is an object of the present invention to use the L3 loop of the Smad proteins 1, 2, 3, 4, 5 or 6 or the C-terminal tail of

Smad proteins 1, 2, 3, 4 or 5 in protein-interaction assays to screen for agents that increase or decrease Smad interactions via these regions.

It is another object of the present invention to provide a method of screening for drugs that interfere with or enhance signaling by TGF-β or other members of the TGF-β family that signal through Smad proteins.

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It is another object of the present invention to provide a screening method that utilizes high specificity peptide-Smad interactions and peptide receptor interactions and is suitable for adaptation to high throughput assays.

In one embodiment of the present invention, there is provided a method of screening for drugs which enhance or inhibit Smad binding to a complementary Smad via the L3 loop region, comprising the steps of: a) producing a synthetic Smad polypeptide encompassing the L3 loop region; b) attaching detectable label onto this polypeptide; c) contacting the synthetic L3 loop polypeptide with a complementary Smad protein immobilized on a solid support; d) measuring the amount of labeled L3 loop polypeptide bound; e) in parallel to steps (c) and (d), conducting these same steps in the presence of a test substance; and f) comparing the amount of L3 loop polypeptide bound in the presence of a test substance with the amount bound in the absence of test substance so as to identify test substances that either increase L3 loop polypeptide binding to the Smad protein or decrease L3 loop polypeptide binding to the Smad protein.

In another embodiment of the present invention, there is provided a method of screening for drugs which enhance or inhibit Smad binding to a complementary Smad via the L3 loop region, comprising the steps of: a) producing a synthetic Smad polypeptide, encompassing the L3 loop region as defined by the crystal structure of the Smad4/DPC4 C-terminal domain; b) producing this polypeptide containing a chemical group that allows immobilization; c) contacting this L3 loop polypeptide with a labeled complementary Smad protein; d) measuring the amount of labeled Smad protein bound to the L3 loop polypeptide; e) in parallel to steps (c) and (d), conducting these same steps in the presence of a test substance; and f) comparing the amount of Smad protein bound in the presence of a test substance with the amount bound in the absence of test substance in order to identify test substances that either increase L3 loop polypeptide binding to the Smad protein or decrease L3 loop polypeptide binding to the Smad protein.

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In yet another embodiment of the present invention, there is provided a method of screening for drugs which enhance or inhibit Smad4 binding to a complementary Smad via the C-terminal phosphorylated tail ("C-tail") of this Smad, comprising the steps of: a) producing a synthetic polypeptide corresponding to the C-terminal tail of a given Smad encompassing the C-terminal tail that follows the H5 alpha-helix as defined by the crystal structure of the Smad4/DPC-terminal domain; b) attaching a detectable label onto this polypeptide; c) contacting this C-tail polypeptide with Smad4 protein immobilized on a solid support; d) measuring the amount of labeled C-tail polypeptide that is bound to Smad4; e) in

parallel to steps (c) and (d), conducting these same steps in the presence of a test substance; and f) comparing the amount of C-tail bound in the presence of a test substance with the amount bound in the absence of the substance in order to identify test substances that either increase C-tail polypeptide binding to the Smad protein or decrease C-tail polypeptide binding to the Smad protein.

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In yet another embodiment of the present invention, there is provided a method of screening for drugs which enhance or inhibit Smad4 binding to a complementary Smad via the Cterminal phosphorylated tail ("C-tail") of this Smad, comprising the steps of: a) producing a synthetic polypeptide corresponding to the C-terminal tail of a given Smad encompassing the C-terminal tail that follows the H5 alpha-helix as defined by the crystal structure Smad4/DPC4 C-terminal of the domain; b) producing polypeptide containing chemical a group that allows immobilization; c) contacting this derivative C-tail polypeptide with the labeled Smad4 protein; d) measuring the amount of labeled Smad4 bound to the C-tail polypeptide; e) in parallel to steps (c) and (d), conducting these same steps, in the presence of a test substance; f) comparing the amount of Smad4 bound in the presence of a test substance, with the amount bound in the absence of test substance in order to identify test substances that either increase Smad4 binding to the C-tail polypeptide decrease Smad4 binding to the C-tail polypeptide.

In yet another embodiment of the present invention, there is provided a method of screening for drugs which enhance or inhibit Smad binding to a receptor of the TGF- $\beta$  family, comprising the steps of: a) producing a synthetic polypeptide

corresponding to the amino acid sequence of a given Smad encompassing the L3 loop region as defined by the crystal structure of the Smad4/DPC4 C-terminal domain; b) attaching a detectable label onto this polypeptide; c) contacting this L3 loop polypeptide with a receptor cytoplasmic domain protein such as a Smadl-derived L3 loop polypeptide with the bone morphogenetic protein receptor cytoplasmic domain, or contacting a Smad2derived L3 loop polypeptide with the TGF-\u03b3 receptor cytoplasmic domain) immobilized on a solid support; d) measuring the amount of labeled L3 loop polypeptide; e) in parallel to steps (c) and (d), conducting these same steps, in the presence of a test substance: and f) comparing the amount of L3 loop polypeptide bound in the presence of a test substance with the amount bound in the absence of test substance in order to identify test substances that either increase L3 loop polypeptide binding to the receptor or decrease L3 loop polypeptide binding to the receptor.

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In yet another embodiment of the present invention, there is provided a method of screening for drugs which enhance or inhibit binding of a Smad N-terminal domain to the C-terminal domain of the same Smad protein, comprising the steps of: a) producing recombinant forms of the N-terminal domain and C-terminal domain polypeptides, with one containing a detectable label and the other containing a moiety allowing immobilization onto a solid support; b) contacting the recombinant N-terminal domain polypeptide with the C-terminal domain polypeptide; c) measuring the amount of labeled domain polypeptide bound; d) in parallel to steps (b) and (c), conducting these same steps, in the presence of a test substance; e) comparing the amount of labeled

polypeptide bound in the presence of a test substance with the amount bound in the absence of a test substance so as to identify test substances that either increase N-terminal domain binding to the C-terminal domain or decrease N-terminal domain binding to the C-terminal domain.

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Smad2 and Smad4 are related tumor suppressors that, in response to TGF-β, form a complex that mediates transcriptional and growth inhibitory responses. The effector function of Smad2 and Smad4 is located in their conserved C-terminal domain (C domain) and inhibited by the presence of their N-terminal domains (N domain). The inhibitory function of the N domain is shown herein to involve a physical interaction with the C domain. preventing the association of Smad2 with Smad4. This inhibitory function is increased in tumor derived forms of Smad2 and 4 that carry a missense mutation in a conserved N domain arginine. The mutant N domains have increased affinity for their respective C domains, inhibit Smad2-Smad4 interaction and prevent TGF-\u03Binduced Smad2-Smad4 association and signaling. Whereas mutations in the C domain disrupt the effector function of the Smads, the N domain arginine mutations inhibit Smad signaling through a gain of autoinhibitory function. Gain of autoinhibitory function provides a novel mechanism of tumor suppressor inactivation.

In the present invention, the crystal structure of the Cterminal domain (CTD) of the Smad4/DPC4 tumor suppressor, was
determined at 2.5 Å resolution and revealed that the
Smad4/DPC4-C-terminal domain forms a crystallographic trimer
through a conserved protein-protein interface to which the

majority of the tumor-derived missense mutations map. These mutations disrupt homo-oligomerization in vitro and in vivo, suggesting that the trimeric assembly of the Smad4/DPC4 C-terminal domain is a critical function in signaling that is targeted by tumorigenic mutations.

Other and further aspects, features, and advantages of the present invention will be apparent from the following description of the presently preferred embodiments of the invention given for the purpose of disclosure.

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### BRIEF DESCRIPTION OF THE DRAWINGS

So that the matter in which the above-recited features, advantages and objects of the invention, as well as others which will become clear, are attained and can be understood in detail, more particular descriptions of the invention briefly summarized above may be had by reference to certain embodiments thereof which are illustrated in the appended drawings. These drawings form a part of the specification. It is to be noted, however, that the appended drawings illustrate preferred embodiments of the invention and therefore are not to be considered limiting in their scope.

Figure 1 shows the structure of the Smad4/DPC4-C-terminal domain consists of a  $\beta$ -sandwich with a three-helix bundle on one end and a collection of three large loops and an  $\alpha$  helix on the other end. Schematic representation of the structure is viewed along the edge of the  $\beta$ -sandwich. The dotted line represents the disordered region between the H3 and H4 helices.

Figures were prepared with the programs MOLSCRIPT<sup>26</sup> and RASTER3D<sup>27</sup>.

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Figure 2 shows the Smad C-terminal domains are conserved and are targeted by tumorigenic highly and developmental mutations. Figure 2A shows the sequence alignment of C-terminal domains of five human Smads 1,8,10 (Smad1, 2, 3, 5 and Smad4/DPC4) and homologues from Drosophila<sup>18</sup> (Mad) and C. elegans<sup>19</sup> (Sma-2, 3, 4), with the Smad4/DPC4-C-terminal domain secondary structure elements indicated below the sequences. Residues that are more than 40% solvent-exposed, have no significant structural roles, and are conserved in at least 6 out of the 9 aligned sequences highlighted in cyan. The 14 missense mutations tabulated above the alignment include tumor-derived Smad4/DPC4 and Smad2 mutations 1,2,4,12,17,28, shaded in yellow, as well as mutations from Drosophila and C. elegans genetic screens 18,19 (developmental mutations, shaded in green). The residues where these mutations occur are in bold face and underlined. Figure 2B shows the mapping of both the missense mutations and the highly conserved and solvent-exposed residues identifies the three-helix bundle and the three-loop/helix region as regions likely to be important for macromolecular recognition events that mediate function. Color coding is the same as in Figure 2A. The amino acid substitution and the residue number from the mutated Smad family members other than Smad4/DPC4 are shown The three structural mutations (Arg441Pro from parentheses. Smad4/DPC4, Leu440Arg and Pro445H from Smad2) are not shown.

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Figure 3 shows that in the crystals, the Smad4/DPC4 C-terminal domain forms a trimer that is targeted by tumorigenic mutations and is likely to be important for Smad function. Figure 3 A shows the three monomers, colored red, blue, and magenta, pack across three identical protein-protein interfaces. derived missense mutations map to five amino acids, shown in yellow, that are involved in inter-molecular contacts. Figure 3B shows a close-up view of a inter-molecular hydrogen bond network involving three residues all of which are mutated in cancer. Coloring is according to Figure 3A. Figure 3C shows a close-up view showing the intermolecular packing of Val370. which is mutated to Asp in cancer, against Phe329, Trp524, and the aliphatic portion of Lys519. The subunit in which Val370 is shown is in space-filling representation, whereas the other subunit is shown as the molecular surface (red mesh). Other intermolecular interactions not mentioned include: van der Waals contacts between the L1 loop of the loop/helix region and the H4 and H5 helices of the three-helix bundle (Tyr353, Val354, and Pro356 wedging in-between His530, Leu533, Leu536, Leu540, and His541); the hydrogen bond networks between Ser368 of the L2 loop and Arg496, Glu526, and His528 of the β-sheet, and between His371 of the L2 loop and Asp332 of the β-sheet. This figure was prepared with the program GRASP<sup>29</sup>. Figure 3D shows that in vivo, tumor-derived trimer interface mutations disrupt both homo- and hetero-oligomerization, whereas a developmental mutation in the L3 loop disrupts only hetero-oligomerization. assay for homo-oligomerization, mammalian COS-1 cells were transiently transfected with Flag-tagged wild-type Smad4/DPC4-

C-terminal domain (WT) and HA-tagged WT or mutant constructs. For hetero-oligomerization, cells were transfected with Flagtagged Smad2 C-terminal domain and HA-tagged Smad4/DPC4 C-WT or mutant terminal domain constructs together with constitutively active TGF-\beta type I receptor construct. The cell lysate was immunoprecipitated with anti-Flag antibody immunoblotted using subsequently anti-HA antibody. Immunoblots indicated that the mutant Smad4/DPC4-C-terminal domains expressed at levels comparable to those of the wild type constructs. Studies with the full-length proteins were performed similarly.

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Figure 4 shows the the size exclusion chromatography indicating that the wild type full-length Smad4/DPC4, but not the tumor-derived mutants, has an apparent molecular weight consistent with that of a trimer. Figure 4A shows that recombinant Smad4/DPC4 protein, purified to near homogeneity, was applied to a Superdex200 gel filtration column where it eluted as an approximately 180 kDa molecule. The fractions were visualized with Coomassie staining. Figure 4B shows that in vitro, tumor-derived trimer interface mutations disrupt homooligomerization, whereas a developmental mutation in the L3 loop has no apparent effect on the homo-oligomerization. Gel filtration fractions of partially purified wild-type and mutant Smad4/DPC4 proteins were analyzed by immunoblots with anti-Smad4/DPC4 antibody.

Figure 5 shows that one face of the disk-like trimer structure may mediate hetero-oligomerization. Figure 5A shows that mutations outside the trimer interface map primarily to L3

loop residues, with the exception of Arg420, which is outside the L3 loop. The face of the trimer shown is opposite to that shown in Figure 3A. Figure 5B shows that a model of hetero-oligomer formation depicting the Smad4/DPC4 and Smad2 C-terminal domain trimers as disks. The approximate positions of the Smad4/DPC4 L3 loops and of the Smad2 sites that get phosphorylated by the receptor kinase<sup>30</sup> are indicated by yellow and green, respectively.

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Figure 6 shows an analysis of Smad4 and Smad2 domain interactions. Figure 6A shows the Smad4 and Smad2 interactions with themselves and each other in a yeast two-hybrid GAD fusions with the indicated portions of Smad4 or Smad2 were tested for interaction with full length or C domains of Smad2 or Smad4 fused to the LexA DNA binding domain. relative strength of the interaction is indicated. Figure 6B shows expression level of HA-tagged Smad4 constructs and Flag-tagged Smad2 constructs was determined by epitope-tag immunoprecipitation from 35S-methionine labeled cells. Figure 6C shows homo-oligomerization of Smad4 or Smad2 C domains. COS cells were transiently transfected with full-length (FL) Smad4 or Smad2 or their C domains (C) (Smad4 amino acids 294-552; Smad2 amino acids 248-467). Versions of the same protein tagged N-terminally with the Flag epitope or C-terminally with the HA epitope were cotransfected. Some cultures were incubated with  $TGF-\beta$  for 1 hour before lysis. Homo-oligomerization was analyzed bv anti-HA immunoblotting of anti-Flag immunoprecipitates. Figure 6D shows hetero-oligomerization of Smad4 and Smad2 deletion constructs. HA-tagged Smad4 deletion

constructs were co-transfected with Flag-tagged Smad2, and Flag-tagged Smad2 deletion constructs were cotransfected with full-length HA-tagged Smad4. TGF-β stimulation (+ lanes) was provided by cotransfection of a constitutively active TGF-β type-I receptor and, additionally, incubation with TGF-β. Smad2-Smad4 interactions were analyzed by anti-Flag immunoblotting of anti-HA immunoprecipitates (top panel) or anti-HA immunoblotting of anti-Flag immunoprecipitates (bottom panel). Figure 6E shows summary of Smad domain contributions to Smad2-Smad4 hetero-oligomerization.

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Figure 7A and Figure 7B show the inhibition of Smad2-Smad4 interaction by N domains. Increasing amounts (1, 2) and 4 mg) of plasmid encoding the Smad4 N domain (amino acids 1-154) or the Smad2 N domain (amino acids 1-185) tagged with the indicated epitopes were cotransfected with the indicated full length or C domain forms of Smad4 and Smad2 into COS cells. association Smad2-Smad4 was determined by anti-Flag immunoprecipitation followed by anti-HA immunoblotting. N and Smad4 expression levels were monitored b y immunoblotting with specific antibodies. Figure 7C shows that N domain expression does not affect C domain homo-oligomerization. Flag-tagged and HA-tagged versions of Smad C domains were cotransfected with the indicated N domain. The levels of Smad4 (Figure 7C) (top panel) or Smad2 (Figure 7C) (bottom panel) homo-oligomers were determined by anti-HA immunoblotting of anti-Flag immunoprecipitates.

Figure 8 shows the effect of N domain deletion and agonist-induced phosphorylation on Smad2-Smad4 interaction.

Figure 8A shows constitutive association of the isolated C domains of Smad4 and Smad2, and further stimulation by TGF-B. Flag-tagged full-length or C domain Smad2 constructs and HAfull-length or C domain Smad4 constructs were cotransfected into COS cells. Cultures were stimulated with TGF-B as indicated. Smad2-Smad4 interactions were analyzed by anti-HA immunoblotting of anti-Flag immunoprecipitates. Figure 8B shows Smad2 C domain phosphorylation in response to TGF-B. Constructs were transiently co-transfected with TBR-I into R-1B/ L17 cells. Transfectants were labeled with <sup>32</sup>P-orthophosphate, and stimulated with TGF-β for 20 minutes as indicated. was immunoprecipitated with anti-Flag antibody and subjected to autoradiography (top panel). Quantitation revealed an 8-fold increase in phosphorylation of Smad2 or Smad2(C) in response to TGF-β. Aliquots of cell lysate were subjected to anti-Flag immunoblotting to control for Smad2 levels (bottom panel). Figure 8C shows the constitutive interaction of Smad4 and Smad2 C domains is independent of TGF-B receptor-mediated phosphorylation. Smad2-Smad4 (full-length or C domain) complex formation was analyzed in the presence or absence of a cotransfected dominant negative T $\beta$ R-I construct [ $T\beta$ R-I(KR)]. Other conditions were as described in Figure 8A.

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Figure 9 shows the biological activity of Smad2 and Smad4 containing tumor-derived N domain mutations. Figure 9A shows that wild type Smad2 induces the paraxial mesoderm marker muscle actin in Xenopus ectodermal explants, whereas Smad2 (R133C) or its N domain alone [Smad2(N)R133C] are

unable to induce it. EF-1a was used as an internal control. Figure 9B shows cotransfection of wild type Smad2 and Smad4 (WT) restores TGF-\u03b3 responsiveness into Smad4-defective MDA-MB468 breast cancer cells, whereas co-transfections including Smad2(R133C) mutant (R), the Smad4(R100T) mutant (R) or both mutants do not. The TGF-\beta responsiveness of these cells was determined using the reporter construct 3TP-lux. Figure 9C shows that overexpression of wild type Smad4 inhibits MDA-MB468 cell proliferation whereas overexpression Smad4(R100C) mutant does not. The proliferative activity of the cells determined by measuring iododeoxyuridine incorporation into DNA. Results are the average  $\pm$  S.D. of triplicate assays.

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Figure 10 shows the gain of autoinhibitory function of Smad4 and Smad2 N domain mutants. Figure 10A shows that mutations inhibit the N domain Smad2-Smad4 interaction. Expression levels of wild type and mutant Smads were determined by epitope-tag immunoprecipitation 35S from methionine labeled, transfected COS cells. HA-tagged wild type (WT) or mutant Smad4 was cotransfected with Flag-tagged Smad4 (for homo-oligomeric interaction) or Flag-tagged Smad2 (for hetero-oligomeric interaction) in COS cells. Likewise, Flag-tagged wild type (WT) or mutant (R133C) Smad2 was cotransfected with HA-tagged Smad2 or HA-tagged Smad4. The indicated cells were stimulated TGF-β. with Homo-oligomerization or heterooligomerization was then determined. Figure 10B shows the N domain interaction with the C domain, and its increase by Flag-tagged N domains indicated at the top were mutations.

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cotransfected with the HA-tagged C domains indicated at the bottom. N domain-C domain interaction was determined by anti-HA immunoblotting of anti-Flag immunoprecipitates. expression levels were monitored by immunoprecipitation from 35S-methionine labeled cells. Figure 10C shows that mutant N domains inhibit the Smad2-Smad4 interaction strongly. Increasing amounts of plasmid DNA encoding wild type (WT) or mutant (R100T) Smad4 N domain (left panel) or wild type (WT) or mutant (R133C) Smad2 N domain (right panel) were cotransfected with Flag-tagged Smad2 C domain and HA-tagged The level of Smad2(C)-Smad4(C) complex was then domain. determined by anti-HA immunoblotting anti-Flag The relative levels of Smad4 N domain immunoprecipitates. expressed in these cells were determined by immunoblotting using anti-Smad polyclonal antibody. The levels of Smad4 or Smad2 N domain protein and Smad2(C)-bound Smad4(C) were quantitated (ImageQuant; Molecular Dynamics) and plotted against each other. Figure 10D shows the N domain inhibition of Smad2-Smad4 signaling function, and its increase by N domain mutations. R-1B/L17 cells were transiently transfected with the indicated constructs and 3TP-lux reporter. Amounts of transfected Smad4 and Smad2 were adjusted so that they would increase luciferase expression synergistically. Increasing amounts (1, 2, 4, and 6 mg) of plasmid DNA encoding wild type or mutant N domains were cotransfected with the Smad4(C)/Smad2(C) combination. Results (luciferase activity in arbitrary units) are the average ± S.D. of triplicate assays.

Figure 11A shows a diagrammatic representation of Smad2, its C domain structure based on Smad4, and amino acid sequence alignment of the Smads starting from the L3 loop to the end. In the C domain structure, arrowheads (1 to 11) represent β-sheets; L1 to L3 represent loops; filled circles represent α-helices. In the amino acid sequence alignment, the conserved amino acids are boxed. The two residues in the L3 loop which are distinct among different Smad groups are highlighted. Figure 11B, Inset, shows the structure of the Smad4 C domain trimer highlighting the L3 loop in each monomer. The close-up shows the L3 loop (yellow) protruding from the core structure. The two group-specific amino acids are indicated in red.

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Figure 12 shows the Smad2 association with the TGF-B receptor does not require its C-tail and is affected by Smad2 phosphorylation. Figure 12A: Smad2-TGF-β receptor interaction was determined by co-transfecting Flag-tagged wild type and mutant Smad2 with wild type TβR-I and TβR-II receptors into COS-1 cells, afffnity-labeling by cross-linking to [125]-TGF-\(\beta\)1, then co-immunoprecipitating Smad2-receptor complex using anti-Flag The immunoprecipitates and aliquots of whole cell antibody. lysates were subjected to SDS-PAGE and autoradiography visualize the Smad2-bound receptors (upper panel) and the total receptor levels (lower panel), respectively. Figure 12B: Smad2 phosphorylation was determined by transfecting Flag-tagged wild type or mutant Smad2 alone (-) or together (+) with TBR-I into R1B/L17 cells. After 48 hours, cells were labeled with [32P]orthophosphate for 2 hours and stimulated with (+) or without (-)

TGF-β1 for 30 minutes. Cell lysates were immunoprecipitated with anti-Flag antibody and the immunoprecipitates analyzed by SDS-PAGE and autoradiography. Figure 12C: Expression of Smad2 constructs was checked by transfecting Flag-tagged Smad2 into COS-1 cells. Forty-eight hours post-transfection, cell lysates were resolved by SDS-PAGE and transferred onto membrane support. Western blotting was carried out using anti-Flag antibody.

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Figure 13 shows the Smad 2 C domain retains the receptor docking ability. COS-1 cells were co-transfected with Flag-tagged wild type or mutant Smad2, wild type (WT) or kinase-defective (KR) TβR-I, and wild type TβR-II, and were affinity-labeled with [125I]TGF-β1. The Smad2-bound and total receptors were resolved by SDS-PAGE and autoradiography as described in Figure 11. Smad2 expression was determined in parallel by western blotting.

Figure 14 shows that the L3 loop specifies Smadreceptor interaction. Figure 14A: Differential binding affinity of Smad1 and Smad2 to the TGF-β receptor complex. Figure 14B:

The L3 loop determines the specificity of Smad-receptor interaction. The interaction between the indicated Smad constructs and the TGF-β receptor complex was assessed as described in Figure 11. Smad construct expression levels as determined by anti-Flag immunoblotting are shown in the bottom panel.

Figure 15 shows the role of the L3 loop and C-tail in the phosphorylation of Smads by the type I receptors. Figure 15A: The L3 loop of Smad2 is necessary for Smad2

phosphorylation in response to TGF-\u03b3. Figure 15B: The L3 loop of Smad2 allows Smad1 to be phosphorylated in response to TGF-B, and the Smad2 C-tail supports optimal phosphorylation. Figure 15C: The L3 loop and C-tail of Smadl allows Smad2 to be phosphorylated in response to BMP. Figure 15D: Smad expression level as determined by anti-Flag immunoblotting. determine inducibility of Smad phosphorylation by TGF-B1 or BMP4, R1B/L17 cells were transfected with the indicated Flagtagged Smad constructs alone (-) or together (+) with either TBR-I BMPR-IB and BMPR-II. Cells were labeled with [32P]orthophosphate for 2 hours and then incubated with (+) or without (-) TGF-β1 or BMP4 for 30 minutes. In parallel transfections, Smad proteins immunoprecipitated from cell lysates using anti-Flag antibody were resolved by SDS-PAGE and transferred onto membrane for western blotting using anti-Flag antibody. Arrow indicates Smad proteins.

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Figure 16A shows the association of the receptor-regulated Smads with Smad4. COS-1 cells transfected with the indicated Flag-tagged Smad1 or 2 constructs, HA-tagged Smad4 and activated TβR-I were treated with TGF-β1 for 1 hour. After Smad complexes were immunoprecipitated using anti-Flag antibody, Smad4 was visualized by western blotting with anti-HA antibody. Figure 16B: COS-1 cells were transfected with wild type Smad2 C-terminally tagged with HA epitope (Smad2-HA) and wild type and mutant Smad2 N-terminally tagged with Flag epitope (F-Smad2) (left panel), or transfected with wild type Smad1 C-terminally tagged with HA epitope (Smad1-HA) and wild

type and mutant Smadl N-terminally tagged with Flag epitope (F-Smadl) (right panel). After 48 h, cells were lysed and immunoprecipitation was carried out with anti-Flag antibody and Smad homomeric complexes were visualized by anti-HA immunoblotting.

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Figure 17 shows the nuclear translocation of Smad1, Smad2 and their derivatives in response to TGF-β1 or BMP2. Figure 17A: Vectors encoding the indicated Flag-tagged Smad contructs alone (Control) or together with either TβR-I(T204D) (TβR-I\*) or BMPR-IB(Q203D) (BMPR-IB\*) were transfected into HepG2 cells. 48 hours post-transfection, cells were incubated with TGF-β1 or BMP2 and immnuofluorescence was visualized with primary mouse anti-Flag antibody and secondary FITC-conjugated goat anti-mouse antibody. Nuclear localization was confirmed with DAPI DNA staining. Figure 17B: Percentage of Smads localized in the nucleus was determined by counting 200 to 300 immunofluorescence-positive cells for each sample.

Figure 18A shows the L45 loop sequences of the TGFβ type I receptor family. Conserved amino acids are boxed. Three groups of functionally related receptors have each a characteristic L45 loop sequence. ALK1 is also known as TSR-1, and ALK2 as ActR-I or Tsk7L. Figure 18B shows R-Smad association with Smad4. Scheme, a TGF-β signal transduction pathway with a type II receptor (II), a type I receptor (I), R-Smad phosphorylation (P), Smad4 (4) and a DNA-binding factor (F). COS1 cells were transfected with Flag-tagged Smad1 or Smad2, HA-tagged Smad4, the indicated wild type (WT) or mutant type I receptors, and the

corresponding type II receptors, TβR-II or BMPR-II. R-Smad binding to Smad4 was determined after incubation with TGF-β or BMP2. Figure 18C shows nuclear translocation of R-Smads induced by wild type and L45 mutant type I receptors. HepG2 cells were transfected with Flag-tagged Smad1 or Smad2, the indicated type I receptors, and their corresponding type II receptors. Cells were incubated with TGF-β1 or BMP2 for 1 h and subjected to anti-Flag immunofluorescence.

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Figure 19 shows that exchanging the L45 loops switches the signaling specificity of TBR-I and BMPR-IB. Figure 19A shows the activation of the TGF-β-responsive reporter 3TPluciferase in TBR-I-defective R1B/L17 cells transfected with wildtype or mutant receptors. Cells were incubated with  $TGF-\beta$  (T) or BMP2 (B), and luciferase activity was determined in triplicate samples. Inset, HA-tagged receptors immunoprecipitated from metabolically labeled cells as controls. Figure 19B shows the activation of the A3-CAT reporter containing activin- and TGF-βresponsive Mix.2 elements. R1B/L17 cells were transfected with Fast1 and receptor constructs. TBR-I transfectants were incubated with TGF-β and BMPR-IB transfectants with BMP2, and CAT activity was determined. Figure 19C shows the activation of the BMP-responsive reporter Xvent.2-luciferase in P19 cells transfected with TBR-II and wild type or mutant TBR-I. Cells were incubated with BMP2 (B) or TGF- $\beta$  (T), and luciferase activity was determined. Figure 19D shows induction of markers of dorsal mesoderm (muscle actin), ventral mesoderm (globin) and

neural tissue (NRP-1) in Xenopus embryos. RNAs encoding the indicated constitutively active receptor forms were injected into the animal pole of two-cell embryos. Expression of muscle actin, globin, NRP-1, and EF-1a (as control) in animal caps from these embryos was determined. Animal caps from uninjected embryos (Control), whole embryos (Embryo) and a sample without reverse transcription (-RT) were included.

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Figure 20A shows the receptor-Smad association in COS-1 cells transfected with the indicated type I receptors, the corresponding type II receptors, and Flag-tagged Smad1(1-454) or Receptors were cross-linked to [125I]TGF-\(\beta\)1 (left Smad2(1-456). panel) or [125I]BMP2 (right panel). Smad-bound receptors were visualized by anti-Flag immunoprecipitation, SDS-PAGE and autoradiography (upper panels). Total cell lysates were analyzed to control for receptor expression (middle panels). Smad by expression was controlled immunoprecipitation from metabolically labeled cells (lower panels). Figure 20B shows Smad phosphorylation determined in L17 cells transfected with Flag-tagged Smads, the indicated type I receptors, and the corresponding type II receptors. Cells were labeled with [32P]phosphate, incubated with TGF-B1 BMP2, or and immunoprecipitated with anti-Flag.

Figure 21A shows the sequence alignment of the MH2 domains of Smad1, 2 and 4, with the Smad4 MH2 domain secondary structure elements indicated below. Identical residues are boxed. Subtype-specific residues map to  $\alpha$ -helix 1 (yellow),  $\alpha$ -helix 2 and its vicinity (purple), the L3 loop (red), and

immediately upstream of the C-terminal receptor phosphorylation motif SS(V/M)S (green). The remaining subtype-specific residues (gray) are scattered in the primary sequence but clustered in the crystal structure near the point of connection to the N-terminal half of the molecule<sup>57</sup>. Figure 21B shows a close-up, lateral view of the Smad4 MH2 crystal structure showing the L3 loop (yellow) with subtype specific residues (red) and the  $\alpha$ -helix 2 (cyan) with subtype-specific residues (magenta). Inset, frontal view of the location of the L3 loop and helix 2 or each MH2 monomer in the crystallographic trimer.

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Figure 22 shows the matching receptor L45 loops and R-Smad L3 loops. Figure 22A shows that L3 loop determines Smad activation by a specific receptor but not Smad interaction with Fastl. COS1 cells were transfected with Flag-tagged Smad constructs, myc-tagged Fastl, and TGF-β receptors or BMP receptors. Cells were incubated with the corresponding receptor ligands, TGF-β1 or BMP4, and Smad association with Fastl was determined. *Ig(H)*, immunoglobulin heavy chain. Figures 22B and C show that TβR-I(LB) rescues the ability of TGF-β to induce Smad2(L1) association with Fastl (B) and activation of the A3-luciferase *Mix.2* reporter (C). R1B/L17 cells transfected with various constructs, as indicated, were incubated with 0.5 nM TGF-β for 20 h, and luciferase activity was measured.

Figure 23 shows the α-helix 2 of Smad2 specifies the
25 interaction with the DNA-binding factor Fast1. Figure 23 A shows the interaction of wild type R-Smads and helix 2 exchange

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mutants with Smad4 and Fast1. HA-tagged Smad4 or myc-tagged Fast1 constructs were cotransfected into COS1 cells with the indicated Flag-tagged forms of Smad1 or Smad2. Transfectants were incubated with TGF- $\beta$  (T) or BMP2 (B) and the associations of R-Smads with Smad4 (upper panel) and with Fast1 (lower panel) were determined. The helix 2 exchange mutants bound Smad4 in response to their agonists, but Smad2(H1) lost the ability to associate with Fast1 whereas Smad2(H1) gained the ability to bind Fast1 in response to BMP. Figure 23B shows the activation of a Mix.2 reporter by wild type R-Smads and helix 2 exchange mutants. L17 cells were cotransfected with the indicated forms of Smad1 or Smad2, Fast1, the A3-luciferase construct, and TGF-B receptors or BMP receptors. Cells were incubated with the corresponding receptor ligands, and luciferase activity Smad2(H1) lost the ability to activate the reporter determined. whereas Smad1(H2) gained the ability to do so in response to BMP. Figure 23C shows Fastl-dependent activation of a GAL4 reporter by Smad1(H2). L17 cells were cotransfected with the indicated forms of Smadl, a Fastl fusion with the DNA binding domain from yeast GAL4, a GAL luciferase reporter, and BMP Cells were incubated with or without BMP2, and receptors. luciferase activity was determined. Figure 23D shows the activation of the Xvent.2-luciferase reporter in P19 cotransfected with TBR-I, TBR-II and the indicated Smad2 constructs. Cells were incubated with or without TGF-\u03b3, and luciferase activity was determined in triplicate samples.

Figure 24 shows the determinants of specificity in  $TGF-\beta$  signal transduction. In the  $TGF-\beta$  or BMP receptor

complexes, the type I receptor recognizes and phosphorylates a specific R-Smad, such as Smad2 in the TGF-B pathway or Smad1 in the BMP pathway<sup>43,52</sup>. The R-Smad then associates with Smad4 and moves into the nucleus. Specific association with the DNAbinding factor Fast1 in the nucleus takes the Smad2-Smad4 complex to specific target genes such as Mix.2, activating their transcription<sup>36,34,49</sup>. Selection of a R-Smad by a receptor is specified by the type I receptor L45 loop and the R-Smad L3 loop, whereas selection of a DNA-binding factor (such as Fastl in the case of Smad2) is specified by the  $\alpha$ -helix 2 of the R-Smad. Exchanging any of these three elements between the TGF-B and BMP receptors or between Smad1 and Smad2 causes a switch in the signaling specificity of these two pathways. Specific activation of other target genes by Smad1 or Smad2 complexes is presumed to involve different DNA-binding partners.

#### DETAILED DESCRIPTION OF THE INVENTION

#### **Definitions**

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In accordance with the present invention, there may be employed conventional molecular biology, microbiology, and recombinant DNA techniques within the skill of the art. Such techniques are explained fully in the literature. See, e.g., Sambrook, Fritsch & Maniatis, "Molecular Cloning: A Laboratory Manual (1982); "DNA Cloning: A Practical Approach," Volumes I and II (D.N. Glover ed. 1985); "Oligonucleotide Synthesis" (M.J. Gait ed. 1984); "Nucleic Acid Hybridization" [B.D. Hames & S.J. Higgins eds. (1985)]; "Transcription and Translation" [B.D. Hames & S.J.

Higgins eds. (1984)]; "Animal Cell Culture" [R.I. Freshney, ed. (1986)]; "Immobilized Cells And Enzymes" [IRL Press, (1986)]; B. Perbal, "A Practical Guide To Molecular Cloning" (1984). Therefore, if appearing herein, the following terms shall have the definitions set out below.

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A "DNA molecule" refers to the polymeric form of deoxyribonucleotides (adenine, guanine, thymine, or cytosine) in its either single stranded form, or a double-stranded helix. This term refers only to the primary and secondary structure of the molecule, and does not limit it to any particular tertiary forms. Thus, this term includes double-stranded DNA found, *inter alia*, in linear DNA molecules (e.g., restriction fragments), viruses, plasmids, and chromosomes. In discussing the structure herein according to the normal convention of giving only the sequence in the 5' to 3' direction along the nontranscribed strand of DNA (i.e., the strand having a sequence homologous to the mRNA).

A "vector" is a replicon, such as plasmid, phage or cosmid, to which another DNA segment may be attached so as to bring about the replication of the attached segment. A "replicon" is any genetic element (e.g., plasmid, chromosome, virus) that functions as an autonomous unit of DNA replication in vivo; i.e., capable of replication under its own control. An "origin of replication" refers to those DNA sequences that participate in DNA synthesis. An "expression control sequence" is a DNA sequence that controls and regulates the transcription and translation of another DNA sequence. A coding sequence is "operably linked" and "under the control" of transcriptional and translational control sequences in a cell when RNA polymerase transcribes the coding sequence into

mRNA, which is then translated into the protein encoded by the coding sequence.

In general, expression vectors containing promoter which facilitate sequences the efficient transcription and translation of the inserted DNA fragment are used in connection with the host. The expression vector typically contains an origin of replication, promoter(s), terminator(s), as well as specific genes which capable are of providing phenotypic selection transformed cells. The transformed hosts can be fermented and cultured according to means known in the art to achieve optimal cell growth.

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A DNA "coding sequence" is a double-stranded DNA sequence which is transcribed and translated into a polypeptide in vivo when placed under the control of appropriate regulatory sequences. The boundaries of the coding sequence are determined by a start codon at the 5' (amino) terminus and a translation stop codon at the 3' (carboxyl) terminus. A coding sequence can include, but is not limited to, prokaryotic sequences, cDNA from eukaryotic mRNA, genomic DNA sequences from eukaryotic (e.g., mammalian) DNA, and even synthetic DNA sequences. polyadenylation signal and transcription termination sequence will usually be located 3' to the coding sequence. A "cDNA" is defined as copy-DNA or complementary-DNA, and is a product of a reverse transcription reaction from an mRNA transcript. "exon" is an expressed sequence transcribed from the gene locus, whereas an "intron" is a non-expressed sequence that is from the gene locus.

Transcriptional and translational control sequences are DNA regulatory sequences, such as promoters. enhancers, polyadenylation signals, terminators, and the like, that provide for the expression of a coding sequence in a host cell. A "cis-element" is a nucleotide sequence, also termed a "consensus sequence" or "motif", that interacts with other proteins which can upregulate or downregulate expression of a specicif gene locus. sequence" can also be included with the coding sequence. sequence encodes a signal peptide, N-terminal to the polypeptide, that communicates to the host cell and directs the polypeptide to the appropriate cellular location. Signal sequences can be found associated with a variety of proteins native to prokaryotes and eukaryotes.

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A "promoter sequence" is a DNA regulatory region capable of binding RNA polymerase in a cell and initiating transcription of a downstream (3' direction) coding sequence. For purposes of defining the present invention, the promoter sequence is bounded at its 3' terminus by the transcription initiation site and extends upstream (5' direction) to include the minimum number of bases or elements necessary to initiate transcription at levels detectable above background. Within the promoter sequence will be found a transcription initiation site, as well as protein binding domains sequences) responsible for the binding of RNA (consensus polymerase. Eukaryotic promoters often, but not always, contain "TATA" boxes and "CAT" boxes. Prokaryotic promoters contain Shine-Dalgarno sequences in addition to the -10 and -35 consensus sequences.

"oligonucleotide" is defined The term as a molecule comprised of two or more deoxyribonucleotides, preferably more than three. Its exact size will depend upon many factors which, in upon the ultimate function and use of the turn, depend oligonucleotide. The term "primer" as used herein refers to an oligonucleotide, whether occurring naturally as in a purified restriction digest or produced synthetically, which is capable of acting as a point of initiation of synthesis when placed under conditions in which synthesis of a primer extension product, which is complementary to a nucleic acid strand, is induced, i.e., in the presence of nucleotides and an inducing agent such as a DNA polymerase and at a suitable temperature and pH. The primer may be either single-stranded or double-stranded and must be sufficiently long to prime the synthesis of the desired extension product in the presence of the inducing agent. The exact length of the primer will depend upon many factors, including temperature, source of primer and use the method. For example, for diagnostic applications, depending on the complexity of the target sequence, the oligonucleotide primer typically contains 15-25 or more nucleotides, although it may contain fewer nucleotides.

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The primers herein are selected to be "substantially" complementary to different strands of a particular target DNA sequence. This means that the primers must be sufficiently complementary to hybridize with their respective strands. Therefore, the primer sequence need not reflect the exact sequence of the template. For example, a non-complementary nucleotide fragment may be attached to the 5' end of the primer, with the remainder of the primer sequence being complementary

to the strand. Alternatively, non-complementary bases or longer sequences can be interspersed into the primer, provided that the primer sequence has sufficient complementarity with the sequence or hybridize therewith and thereby form the template for the synthesis of the extension product.

As used herein, the terms "restriction endonucleases" and "restriction enzymes" refer to enzymes which cut double-stranded DNA at or near a specific nucleotide sequence.

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"Recombinant DNA technology" refers to techniques for uniting two heterologous DNA molecules, usually as a result of in vitro ligation of DNAs from different organisms. Recombinant DNA molecules are commonly produced by experiments in genetic engineering. Synonymous terms include "gene splicing", "molecular cloning" and "genetic engineering". The product of these manipulations results in a "recombinant" or "recombinant molecule".

A cell has been "transformed" or "transfected" with exogenous or heterologous DNA when such DNA has been introduced inside the cell. The transforming DNA may or may not be integrated (covalently linked) into the genome of the cell. In prokaryotes, yeast, and mammalian cells for example, transforming DNA may be maintained on an episomal element such as a vector or plasmid. With respect to eukaryotic cells, a stably transformed cell is one in which the transforming DNA has become integrated into a chromosome so that it is inherited by daughter cells through chromosome replication. This stability is demonstrated by the ability of the eukaryotic cell to establish cell lines or clones comprised of a population of daughter cells

containing the transforming DNA. A "clone" is a population of cells derived from a single cell or ancestor by mitosis. A "cell line" is a clone of a primary cell that is capable of stable growth in vitro for many generations. An organism, such as a plant or animal, that has been transformed with exogenous DNA is termed "transgenic".

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As used herein, the term "host" is meant to include not only prokaryotes but also eukaryotes such as yeast, plant and animal cells. A recombinant DNA molecule or gene can be used to transform a host using any of the techniques commonly known to those of ordinary skill in the art. One preferred embodiment is the use of a vectors containing coding sequences for a gene for purposes of prokaryotic transformation. Prokaryotic hosts may include E. coli, S. tymphimurium, Serratia marcescens and Bacillus subtilis. Eukaryotic hosts include yeasts such as Pichia pastoris, mammalian cells and insect cells, and more preferentially, plant cells, such as Arabidopsis thaliana and Tobaccum nicotiana.

Two DNA sequences are "substantially homologous" when at least about 75% (preferably at least about 80%, and most preferably at least about 90% or 95%) of the nucleotides match over the defined length of the DNA sequences. Sequences that are substantially homologous can be identified by comparing the sequences using standard software available in sequence data banks, or in a Southern hybridization experiment under, for example, stringent conditions as defined for that particular system. Defining appropriate hybridization conditions is within the skill of the art. See, e.g., Maniatis et al., supra; DNA Cloning, Vols. I & II, supra; Nucleic Acid Hybridization, supra.

A "heterologous' region of the DNA construct is an identifiable segment of DNA within a larger DNA molecule that is not found in association with the larger molecule in nature. Thus, when the heterologous region encodes a mammalian gene, the gene will usually be flanked by DNA that does not flank the mammalian genomic DNA in the genome of the source organism. In another example, the coding sequence is a construct where the coding sequence itself is not found in nature (e.g., a cDNA where the genomic coding sequence contains introns, or synthetic sequences having codons different than the native gene). Allelic variations or naturally-occurring mutational events do not give rise to a heterologous region of DNA as defined herein.

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In addition, the invention may also include fragments (e.g., antigenic fragments or enzymatically functional fragments) of a gene. As used herein, "fragment," as applied to a polypeptide, will ordinarily be at least 10 residues, more typically at least 20 residues, and preferably at least 30 (e.g., 50) residues in length, but less than the entire, intact sequence. Fragments can be generated by methods known to those skilled in the art, e.g., by enzymatic digestion of naturally occurring or recombinant proteins, by recombinant DNA techniques using an expression vector that encodes a defined fragment, or by chemical synthesis. The ability of a candidate fragment to exhibit a characteristic (e.g., binding to a specific antibody, or exhibiting partial enzymatic or catalytic activity) can be assessed by methods described herein. Purified fragments or antigenic fragments can be used to generate new regulatory enzymes using multiple functional fragments from

different enzymes, as well as to generate antibodies, by employing standard protocols known to those skilled in the art.

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A standard Northern blot assay can be used to ascertain the relative amounts of mRNA in a cell or tissue obtained from plant or other transgenic tissue, in accordance with conventional Northern hybridization techniques known to those persons of ordinary skill in the art. Alternatively, a standard Southern blot assay may be used to confirm the presence and the copy number of the gene in transgenic systems, in accordance with conventional Southern hybridization techniques known to those of ordinary skill in the art. Both the Northern blot and Southern blot use a hybridization probe, e.g. radiolabelled cDNA, either containing the full-length, single stranded DNA or a fragment of that DNA sequence at least 20 (preferably at least 30, more preferably at least 50, and most preferably at least 100 consecutive nucleotides in length). The DNA hybridization probe can be labelled by any of the many different methods known to those skilled in this art.

The labels most commonly employed for these studies are radioactive elements, enzymes, chemicals which fluoresce when exposed to untraviolet light, and others. A number of fluorescent materials are known and can be utilized as labels. These include, for example, fluorescein, rhodamine, auramine, Texas Red, AMCA blue and Lucifer Yellow. A particular detecting material is antirabbit antibody prepared in goats and conjugated with fluorescein through an isothiocyanate. Proteins can also be labeled with a radioactive element or with an enzyme. The radioactive label can be detected by any of the currently available counting procedures.

The preferred isotope may be selected from <sup>3</sup>H, <sup>14</sup>C, <sup>32</sup>P, <sup>35</sup>S, <sup>36</sup>Cl, <sup>51</sup>Cr, <sup>57</sup>Co, <sup>58</sup>Co, <sup>59</sup>Fe, <sup>90</sup>Y, <sup>125</sup>I, <sup>131</sup>I, and <sup>186</sup>Re.

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Enzyme labels are likewise useful, and can be detected by any of the presently utilized colorimetric, spectrophotometric, fluorospectrophotometric, amperometric or gasometric techniques. The enzyme is conjugated to the selected particle by reaction with molecules bridging such as carbodiimides. diisocvanates. glutaraldehyde and the like. Many enzymes which can be used in these procedures are known and can be utilized. The preferred peroxidase, β-glucuronidase, β-D-glucosidase, galactosidase, urease, glucose oxidase plus peroxidase and alkaline phosphatase. U.S. Patent Nos. 3,654,090, 3,850,752, and 4,016,043 are referred to by way of example for their disclosure of alternate labeling material and methods.

The following specific definitions are given for the purposes of describing the art to which the present invention pertains specifically and distinctly. Any terms not specifically defined herein have the meaning generally known in this art.

As used herein, the term "Smad4/DPC4 and Smad2" shall refer to two related cytoplasmic proteins of known amino acid sequence that mediate the effects of TGF-β and that form a complex with each other in response to stimulation with TGF-β.

As used herein, the term "receptor-regulated Smad polypeptide" or "receptor-regulated Smad protein" shall refer to a minimum of seven cytoplasmic proteins of known amino acid sequence that mediate the effects of TGF- $\beta$  and are contacted by the TGF- $\beta$  receptors.

As used herein, the term "TGFβ/activin/bone morphogenetic protein (BMP)-2/4 cytokine superfamily" shall refer to a family of related polypeptide growth factors of known amino acid sequence.

As used herein, the term "protein-interaction assay" shall refer to an assay that measures, or depends upon, the specific association of one protein with another. The association may occur between these proteins in solution or inside cells.

As used herein, the term "effector function" shall refer to the ability to generate or activate specific cellular responses.

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As used herein, the term "autoinhibitory function" shall refer to the ability of one portion of the Smad protein to inhibit or repress the effector function of another portion of the same protein.

As used herein, the term "tumor-derived missense mutation" shall refer to an amino acid change originated by a single base mutation found in a human tumor sample.

As used herein, the term "homo-oligomerization and hetero-oligomerization" shall refer to the process and ability of a Smad protein to associate with itself, i.e., homo-oligomerize, or to associate with another Smad protein, i.e., hetero-oligomerize.

As used herein, the term "L3 loop region" shall refer to a region in the carboxy-terminal domain of Smad proteins whose length and boundries are defined by the crystal structure of the Smad4/DPC4 C-terminal domain and is expressed on the surface of this domain. Mutation of the L3 loop region prevents Smad hetero-oligomerization and receptor association without preventing Smad homo-oligomerization.

As used herein, the term "loop/helix region" shall refer to a Smad C-terminal domain region defined by the crystal structure of Smad4/DPC4 and involved in Smad homooligomerization by interaction with the three helix bundle.

As used herein, the term "L45 loop region" shall refer to a region of known amino acid sequence in the TGF-β receptors that is required for these receptors to contact and recognize receptor-regulated Smads.

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As used herein, the term "α-helix 2 of the MH2 domain" shall refer to a region of known amino acid sequence in the Smad proteins that is required by these proteins to contact and recognize DNA binding factors.

As used herein, solid support shall refer to a matrix to which a protein or nucleic acid molecule may be attached, for example, by covalent means. For purposes of example, a solid support may comprise matrices consisting of agarose, sepharose, polyacrylamide, nitrocellulose, polystyrene and PVDF.

As used herein, the term " $\beta$ -sandwich" shall refer to the core structure of the C-terminal domain of the Smad protein as defined by the crystal structure of Smad4.

As used herein, the term "three-helix bundle" shall refer to a region of the Smad protein C-terminal domain defined by the crystal structure of Smad4/DPC4 and is involved in Smadhomo-oligomerization by interaction with the loop/helix region.

As used herein, the term "invariant" shall refer to an amino acid residue that remains the same in all Smad proteins at a given position in their amino acid sequence.

Mediation of growth inhibitory responses (such as cell cycle arrest, terminal differentiation and/or apoptosis) and the induction of extracellular matrix proteins (such as collagens, fibronectin, proteoglycan) are important biochemical events. In cancer, mutation in the Smad2 or Smad4 receptor is known to inactivate certain biochemical pathways which deprive the cell of growth inhibitory mechanisms. In fibrotic disorders of the kidney, liver and lung, the TGFβ-Smad pathway is hyperactive. Thus, agents which enhance the function of the pathway would be beneficial in the treatment of cancer whereas agents that inhibit the pathway would be beneficial in the treatment of fibrosis. The present invention discloses that such manipulation of the TGFB-Smad pathway is possible by focusing on the interaction between specific receptor-activated Smads. These Smads interact with the receptor through specific contacts as described in detail below. Upon phosphorylation by the receptor, these Smads dissociate and form a complex with Smad4. Smad4 itself is not a receptor substrate but its association with Smads 1, 2 or others is essential for the transcriptional activity of these complexes.

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The present invention discloses which regions of the Smad protein are involved in the Smadl-receptor or Smad2-receptor interaction and which regions of the Smad protein are involved in the Smadl-Smad4 interation. Discrete differences in the amino acid sequence of specific regions within these domains dictate whether a Smad protein will interact with a given TFG\$\beta\$ family receptor. Structures within this domain also mediate the crucial interaction between Smad4 and Smads 1, 2, 3 or 5.

The present invention discloses that the L3 loop region of the Smad4 protein is exposed on the surface of Smad4 and is conserved in all other Smads. However, certain amino acid residues within this loop vary in each Smad. Furthermore, of several mutations previously identified in inactive alleles of Smad, three fall in the L3 loop of these Smads. The L3 loop mutations do not affect the homotrimeric contacts between the Smad subunits but do eliminate the Smad4 interaction with other Smads. Thus, the L3 loop is the structural motif that mediates Smad4 contact with Smads 1, 2, 3 and 5. The L3 loop is also required for Smad 1, 2, 3 or 5 interaction with the receptor. As discussed below, the crystal structure of Smad4 reveals how the C-terminal tail containing the last few amino acids of a Smad emerges from the globular structure. In Smads 1, 2, 3 and 5, this tail contains the receptor phosphorylation sites. The crystal structure of Smad4 illustrates exactly where this tail starts.

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The present invention is directed to the use of specfic L3 loop peptides or C-tail peptides as ligands for recombinant forms of other Smads, e.g., the Smad1 L3 loop as a ligand of Smad4, the Smad4 L3 loop as a ligand of Smad1, or the L3 loop as a ligand of type 1 receptors. Using the loop region alone as a ligand affords greater specificity in the assays. This assay can be used to screen for drugs which either enhance or inhibit Smad binding.

Thus, the present invention provides a method of testing compounds, comprising the steps of: a) providing (i) a Smad4 polypeptide comprising the L3 loop region, (ii) a complementary Smad polypeptide, and (iii) a compound to be

(b) contacting said Smad4 polypeptide tested; with said complementary Smad polypeptide under conditions where binding can take place, wherein said contacting is performed in the presence and absence of said compound; and c) detecting an increase or decrease in binding of said Smad4 polypeptide to said complementary Smad polypeptide in the presence of said compound. Preferably, the complementary Smad polypeptide is selected from the group consisting of Smad1, Smad2, Smad3, Smad5 and Smad8.

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The present invention also provides a method of 10 testing compounds, comprising the steps of: a) providing (i) two Smad polypeptides from the same Smad family comprising the Cterminal domains of each, and (ii) a compound to be tested; b) contacting said Smad polypeptides under conditions binding can take place, wherein said contacting is performed in the presence and absence of said compound; and c) detecting an increase or decrease in binding of said Smad polypeptides to each other in the presence of said compound. Preferably, the families of Smad polypeptides are selected from the group consisting of 20 Smad1, Smad2, Smad3, Smad4, Smad5, Smad6, Smad7 and Smad8.

The present invention also provides a method of testing compounds, comprising the steps of: a) providing (i) a Smad polypeptide comprising the C-terminal domain, polypeptide comprising the L45 loop of the kinase domain corresponding to a receptor of the TGF-\_ or BMP family, and (iii) a test compound; b) contacting said Smad polypeptide with said receptor polypeptide under conditions where phosphorylation can take place, wherein said contacting is performed in the presence

and absence of said compound; and c) detecting an increase or decrease in the phosphorylation of said Smad polypeptide in the presence of said compound. Preferably, the Smad polypeptide is selected from the group consisting of Smad1, Smad2, Smad3, Smad5 and Smad8.

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The present invention also provides a method of testing compounds, comprising the steps of: a) providing (i) a Smad polypeptide comprising the (-helix 2 of the MH2 domain, (ii) a DNA binding polypeptide, and (iii) a compound to be tested; b) contacting said Smad polypeptide with said DNA binding polypeptide under conditions where binding can take place. wherein said contacting is performed in the presence and absence of said compound; and c) detecting whether there is an increase in binding of said Smad polypeptide to said DNA binding polypeptide in the presence of said compound. Preferably. the polypeptide is selected from the group consisting of Smad1, Smad2, Smad3, Smad4, Smad5 and Smad8. Preferably, the DNA binding polypeptide is selected from the group consisting of FAST1 and homologues of FAST1.

The present invention also provides a method of testing compounds, comprising the steps of: a) providing (i) two Smad polypeptides comprising the C-terminus of each, (ii) a Smad polypeptide comprising the N-terminal domain, and (iii) a compound to be tested; b) contacting said Smad C-terminus polypeptides in the presence of said Smad N-terminal domain under conditions where binding can take place, wherein said contacting is performed in the presence and absence of said compound; and c) detecting whether there is an increase or

decrease in binding of said Smad C-terminus domains in the presence of said compound due to inhibition of the autoinhibitory function of the N-terminal domain by said compound. Preferably, the Smad polypeptide is selected from the group consisting of Smad1, Smad2, Smad3, Smad4, Smad5 and Smad8.

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The present invention also provides a method of testing compounds, comprising the steps of: a) providing (i) a Smad polypeptide comprising the C-terminal domain, (ii) a polypeptide comprising the L45 loop of the kinase domain corresponding to a receptor of the TGF-\_ or BMP family, and (iii) a test compound; b) contacting said Smad polypeptide with said receptor polypeptide under conditions where binding can take place, wherein said contacting is performed in the presence and absence of said compound; and c) detecting an increase or decrease in the binding of said Smad polypeptide to said kinase domain in the presence of said compound. Preferably, the Smad polypeptide is selected from the group consisting of Smad1, Smad2, Smad3, Smad5 and Smad8.

The compounds tested in the methods of the present 20 invention may be used to treat a variety of ailments. Representative ailments include pancreatic cancer, breast cancer, ovarian cancer, colon cancer, esophageal cancer, head and neck cancers, fibrosis of the kidney, fibrosis of the liver, fibrosis of the lung, Alzheimer's disease, memory loss, inflammation, wound 25 healing, bone growth, immunoregulation, blood cell formation and atherosclerosis.

A person having ordinary skill in this art would readily recognize that a variety of detection techniques may be

utilized in the methods of the present invention. Representative detection techniques include solid support immobilization of one or the other polypeptides, labeling of one or the other polypeptides, scintillation proximity, homogeneous time resolved fluorescence, fluorescence resonance energy transfer and fluorescence polarization.

The following examples are given for the purpose of illustrating various embodiments of the invention and are not meant to limit the present invention in any fashion.

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## EXAMPLE 1

## Protein expression and purification

Recombinant Smad4/DPC4-C-terminal domain, corresponding to residues 319-552, was overexpressed at room temperature in *Escherichia coli* using a pET vector (Novagen). The Smad4/DPC4-C-terminal domain in the soluble fraction of the *E coli* lysate was partially purified on a Q-Sepharose column, was concentrated by ultrafiltration and was further purified by gel filtration chromatography (Superdex75 column) and by anion-exchange chromatography (Source 15Q column).

### EXAMPLE 2

### Crystallization

Initial crystals were grown at 4°C by the hanging-drop vapour-diffusion method, by mixing the 10-15 mg/ml protein solution with an equal volume of the reservoir solution containing 100 mM MES, 25% monomethylether PEG5000 (MPEG5000), and 200 mM (NH<sub>4</sub>)<sub>2</sub>SO<sub>4</sub> (pH6.5). Crystals suitable for diffraction studies

were grown using streak-seeding and macroseeding methods<sup>20</sup>. The crystals form in the cubic space group F4132 with a = b = c = 199.6 Å, and contain one molecule in the asymmetric unit.

# EXAMPLE 3

## Data collection and processing

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Diffraction data were collected using an R-AXISIIC imaging plate detector mounted on a Rigaku 200HB generator. Nativel and derivative data were collected at 8°C, and native2 data were collected at -170°C with a crystal flash frozen in a buffer containing 20% glycerol and 25% MPEG5000. Heavy-atom soaks were performed in 50 mM HEPES, 25% MPEG, 160 mM (NH<sub>4</sub>)<sub>2</sub>SO<sub>4</sub>, 100 mM NaCl, pH 6.1, containing one of the following heavy-atom solutions: 1.2 mM thimerosal for 12 hours, 3.0 mM (CH<sub>3</sub>)<sub>3</sub>PbCOOCH<sub>3</sub> for 3 days, and 2.0 mM uranyl acetate for 19 hours.

#### EXAMPLE 4

## MIR analysis, model building and refinement

The heavy atom sites of the thimerosal derivative were determined by direct methods with the program SHELXS-90<sup>21</sup>, and the heavy atom sites of the other derivatives were identified by difference fourier methods. Initial MIR phases calculated with the program MLPHARE<sup>22</sup> had a mean figure of merit of 0.62 to 3.2 Å, and they were improved with solvent flattening and histogram matching with the program SQUASH<sup>23</sup>. The MIR electron density maps had continuous electron density for the majority of the Smad4/DPC4-C-terminal domain

polypeptide, with the exception of a 34 amino acid region between helices H3 and H4. A model was built into MIR electron density maps with the program O<sup>24</sup>, it was refined by simulated annealing with the program X-PLOR<sup>25</sup>, and it was checked by calculating X-PLOR omit maps in which 5-7% of the structure was deleted in each calculation and simulated annealing was used to reduce model bias. The refined model contains residues 319-543 of human Smad4/DPC4 and 129 water molecules. Residues 544-552 at the C-terminus, and residues 457-491 between helices H3 and H4 have no electron density in the maps and it is likely that these regions were disordered in the crystals.

## **EXAMPLE 5**

#### In vivo oligomerization assays

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The full-length Smad4/DPC4 and Smad2, Smad4/DPC4-C-terminal domain encoding amino acids 294-552, Smad2-C-terminal domain encoding amino acids 248-467 were subcloned into the mammalian expression vector pCMV5. Smad4/DPC4 point mutations were generated by a polymerase chain reaction (PCR)-based method and were confirmed by DNA sequencing. Mammalian COS-1 cells were transiently transfected with the indicated FLAG- and HA-tagged constructs by the DEAEdextran method. Two days after transfection, cells were incubated with 200 pM TGF-\beta1 for one hour. Cells were lysed and subjected to immunoprecipitation followed by immunoblot as described9. wash buffers contained 150 mM NaCl for all immunoprecipitation experiments except for the homooligomerization assays of the full-length wild-type Smad4/DPC4

and point mutants, where 250 mM NaCl was employed to better differentiate the WT and mutant activities.

## EXAMPLE 6

### 5 In vitro oligomerization assays

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The full-length Smad4/DPC4 proteins, both wild-type and point mutants, were overexpressed at room temperature in E.coli using a pET vector (Novagen). Smad4/DPC4 protein in the soluble fraction of the E. coli lysate was partially purified by ion exchange chromatography (Q-Sepharose) and was applied to a gel filtration column (Superdex200) in 50 mM Tris, 200 mM NaCl, 5 mM DTT, pH 8.0. Aliquots from the fractions corresponding to molecular weight standards between 440 kDa and 25 kDa were taken for immunoblots with a rabbit polyclonal antibody raised against the Smad4/DPC4-C-terminal domain. The results were visualized with the ECL Western analysis and detection system (Amersham). In addition, the WT full-length Smad4/DPC4 was also cloned as a GST-fusion protein and purified to near homogeneity over a glutathione column. The fusion protein was then cleaved with Thrombin and the Smad4/DPC4 protein was futher purified by anion-exchange chromatography (Source 15Q column).

To help understand how the Smad C-terminal domain functions in mediating TGF $\beta$  signaling and how its mutation in cancer inactivates the pathway, the crystal structure of the 234 amino acid Smad4/DPC4-C-terminal domain (residues 319 to 552) at 2.5 Å resolution (TABLE 1) was determined. The structure consists of a  $\beta$ -sandwich with twisted antiparallel  $\beta$ -sheets of five

and six strands each (Figure 1). One end of the  $\beta$ -sandwich is capped by a three- $\alpha$ -helix bundle (H3, H4, and H5 helices) that extends over the plane of the six-stranded  $\beta$ -sheet, at a roughly perpendicular angle; the other end of the  $\beta$ -sandwich is capped by a group of three large loops and an  $\alpha$ -helix (L1, L2, L3 loops, and H1 helix; Figure 1).

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TABLE 1

Statistics from the crystallographic analysis

2	Data set	Native1 (8 °C)	-	Native2 (-170 °C)	Thimerosal	sal	UO <sub>2</sub> (OAc) <sub>2</sub>	PbOAc
	Resolution (Å) Observations	3.0		2.5 39125	3.0		3.0	3.2
0	Unique reflections	7189		1496	7073		6765	5759
	Data coverage (%)	96.8		96.5	96.8		92.8	94.6
	R <sub>sym</sub> (%)	6.5		7.1	4.8		8.5	10.0
v	MIR analysis (20.0-3.2 Å):	(			0		7	600
<b>ว</b>	Mean isomorphous umenence	92			00		- 0	42.0
49	Phasing power Refinement statistics:				2.54		1.38	7.02
)							RMSD	D
	Resolution Reflections	Protein	Waters	R-factor	R-free	ponds	angles	B-factor
0	$(\dot{A})$ $(IFI > 2\sigma)$	atoms	atoms	(%)	(%)	<b>(Å</b> )	(0)	$(A^2)$
		1522	129	20.9	28.6	0.010	1.66	3.29
		;	,					

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Rsym =  $\Sigma_h \Sigma_i II_h + I_h I \Sigma_h \Sigma_i I_{h,i}$  for the intensity (1) of i observations of reflection h. Mean isomorphous difference = =  $[(F_{H(calc)}^2)/(F_{PH(obs)}^2-F_{PH(calc)})^2]^{1/2}$ . Figure of merit =  $|F(hk)_{best}|/F(hk)$ . R-factor =  $\Sigma |F_{obs}^2-F_{calc}|/\Sigma|F_{obs}^1$ , where Fobs and Foalc are the observed and calculated structure factors, respectively. R-Free = R-factor calculated using 5% of the reflection data chosen randomly and omitted from the start of refinement. RMSD: root mean  $\Sigma$ IFpH-FpI/ $\Sigma$ FpH, where FpH and Fp are the derivative and native structure factors, respectively. Phasing power square deviations from ideal geometry and root mean square variation in the B-factor of bonded atoms.

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To simplify the presentation, the three large loops and  $\alpha$ -helix, as well as portions of  $\beta$ -strands in their immediate vicinity are referred to collectively as the loop/helix region. The three  $\alpha$ -helices of the bundle pack in an up-down-up orientation primarily through leucine residues. In-between the H3 and H4 helices, a 34 amino acid sequence that is rich in Ala (39%), Gly and Pro residues and is present only in Smad4/DPC4 and its *C. elegans* homologue Sma-4, is disordered in the crystals (residues 457 to 491). In the loop/helix region, the L1, L2, and L3 loops of 7, 9, and 18 residues, respectively, and the H1 helix are mostly polar and pack through extended hydrogen bond networks. These hydrogen bonds are likely to contribute to the rigid structure of this region that is suggested by the well-defined electron density.

Smad proteins are highly conserved within the family and across species, with Smad4/DPC4 and its *C. elegans* homologue, Sma-4, representing a somewhat divergent subtype which still retains about 40% identity with other family members<sup>5-7</sup> (Figure 2A). Many of the conserved residues have structural roles. These include the hydrophobic residues that make up the hydrophobic core of the β-sandwich and of the three-helix bundle, as well as many of the polar residues that form the hydrogen bond networks important for the structure of the loop/helix region. Examples of the latter group are the invariant Arg372 and Arg380 residues from the H1 helix making 4 and 3 charge stabilized hydrogen bonds, respectively. Many other highly conserved residues are solvent-exposed and have

no apparent structure-stabilizing roles. They are thus candidates for functional residues that may mediate macromolecular interactions important for the function of Smad proteins. The structure reveals that these candidate functional residues, which are highlighted in Figure 2B, show a strong tendency to cluster at the loop/helix region and the three-helix bundle.

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Besides sequence conservation, another indication that the loop/helix region and the three-helix bundle are functionally important comes from an analysis of the 9 tumor-derived missense mutations, some observed multiple times, in the C-terminal domains of the Smad4/DPC4 and Smad2 tumor suppressors. Excluding three mutations that map to structural residues, 5 of the 6 tumor-derived missense mutations map to either the loop/helix region or to the three helix bundle: the Smad4/DPC4 mutations Asp351His<sup>2</sup>, Arg361Cys<sup>17</sup>, and Val370Asp<sup>17</sup> map to the loop/helix region, whereas the Smad4/DPC4 mutation Asp493His1 and the Smad2 mutation Asp450Glu<sup>12</sup> (corresponding to Asp537 of Smad4/DPC4) map to the three-helix bundle. These mutations may deprive the C-terminal domain of critical intermolecular contacts.

The one mutation that does not map to either region is Arg420His from Smad4/DPC4, which instead maps to the side of the  $\beta$ -sandwich (H2 helix), a region that is not as well conserved. The remaining three mutations map to structural residues: the Smad2 Leu440Arg mutation (corresponding to Ile527 of Smad4/DPC4) in the hydrophobic core of the  $\beta$ -sandwich likely disrupts the packing in the hydrophobic core; the Smad4/DPC4 Arg441Pro mutation at the

three-helix bundle likely disrupts the H3 helix because of the introduction of a proline in the midst of the helix; and the Smad2 Pro445His mutation (corresponding to Ala532 in Smad4/DPC4), also at the three-helix bundle, likely disrupts the packing between the three-helix bundle and the  $\beta$ -sandwich as there is little space for the bigger histidine side chain in this portion of the hydrophobic core.

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Additional support for the functional significance of the loop/helix region is provided by mutations in *Drosophila* and *C elegans* that produce null or severe developmental phenotypes 18,19. These developmental mutations map to Gly508 (*Drosophila* Mad, *C elegans* Sma-2), Gly510 (Sma-3), and Glu520 (Mad) of the L3 loop in the loop/helix region (Figure 2). Thus, the locations of conserved, solvent-exposed residues and the locations of mutations derived from tumors or from *Drosophila* and *C. elegans* genetic screens, taken together, point to the loop/helix region and the three-helix bundle as playing a critical role in mediating Smad activities.

Because the Smad C-terminal domains can mediate most of the biological effects of the full-length proteins, the Smad4/DPC4-C-terminal domain was tested for the homo-oligomerization activity. Initial co-immunoprecipitation experiments using extracts from COS cells transfected with differentially tagged Smad4/DPC4-C-terminal domain constructs showed that the Smad4/DPC4-C-terminal domain retained the ability to form homo-oligomers when overexpressed (Figure 3D), suggesting that the C-terminal domain may contain a primary homo-oligomerization activity. However, the full-length Smad4/DPC4 homo-oligomers are more stable than the Smad4/DPC4-

C-terminal domain homo-oligomers in vivo, suggesting that residues N-terminal to the Smad4/DPC4-C-terminal domain are likely to contribute to homo-oligomerization.

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To further investigate the homo-oligomerization activity of the Smad4/DPC4-C-terminal domain. the packing of the Smad4/DPC4-C-terminal domain molecules in the crystals was examined and a crystallographic trimer that formed through three identical, extended protein-protein interfaces, burying a total of 4800 Å<sup>2</sup> of surface area was identified (Figure 3A). Each interface forms through the interactions of the highly conserved regions of the Smad4/DPC4-C-terminal domain that contain the majority of the candidate functional residues: the loop/helix region of one subunit packs extensively with the three-helix bundle from another subunit. while making a few additional contacts to residues from the Bsandwich (Figure 3A). The only portion of the loop/helix region that does not participate in this interface is the L3 loop.

The trimer interface includes the majority of the conserved residues and the tumor-derived non-structural missense mutations (five out of six). Most noteworthy is an extended intermolecular hydrogen bond network involving, from one subunit, the Arg361 and Asp351 side chains and two backbone amide groups of the loop/helix region, and from another subunit, the Asp537 side chain of the three-helix bundle (Figure 3B). The Asp351, Arg361, and Asp537 residues are essentially invariant, with the exception of a conservative Arg to Lys substitution in Sma-2 (Figure 2A), and all three are mutated in cancer (Figure 2). The Asp351His and

Arg361Cys mutations have been isolated from Smad4/DPC4 in ovarian<sup>2</sup> and colon cancer<sup>17</sup>, respectively, and the Asp450Glu mutation, corresponding to Asp537 of Smad4/DPC4, has been isolated from Smad2 in colon cancer<sup>12</sup>. Each of these mutations is certain to disrupt this intricate hydrogen bond network at the interface. Also noteworthy are the intermolecular van der Waals contacts between Val370 on the L2 loop of the loop/helix and the Trp524, Phe329, and the aliphatic portion of the Lys519 side chain on the  $\beta$ -sheet at the base of the three-helix bundle (Figure 3C).

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The two aromatic residues are also essentially invariant, with the exception of a conservative Tyr to Phe substitution in Smad4/DPC4 (Figure 2A). Furthermore, Val370 is found mutated to Asp in colon cancer<sup>17</sup>. The introduction of a charged amino acid into a hydrophobic portion of the interface should be effective in destabilizing the trimer interface. Finally, the Smad4/DPC4 Asp493His mutation from pancreatic cancer<sup>1</sup> also maps to the trimer interface (Figure 3A) and would interfere with the electrostatic packing of Asp493 of one subunit with Arg496 and Arg497 of another subunit at the trimer interface. However, in the crystals, Asp493 is near the disordered region of the H4 helix and its interactions with the arginines are not well defined.

Many of the other trimer-interface contacts are also conserved in the Smad family (Figure 3C), indicating that other Smad-C-terminal domains may form a similar trimeric structure. On the other hand, not all residues in the Smad4/DPC4-C-terminal domain trimer interface are conserved in all Smads, and it is likely

that those that differ may contribute to subtype specificity. An example of this is an intermolecular hydrogen bond contact between His371 and Asp332. This pair is conserved in the *C. elegans* Smad4/DPC4 homologue, Sma4, whereas it is an invariant Asn-Asn pair in the pathway-restricted Smads (Figure 2).

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If the trimeric Smad4/DPC4-C-terminal domain assembly observed in the crystals is part of the homo-oligomer observed in vivo, then mutations at residues that make intermolecular contacts at the interface, and in particular the tumor-derived mutations discussed earlier, should disrupt or reduce homo-oligomerization in Figure 3D shows the results of co-immunoprecipitation vivo. experiments using extracts from COS cells transfected differentially tagged mutant Smad4/DPC4 molecules. All four of the tumorigenic mutations at residues that play important roles in the trimer interface, Asp351, Arg361, Val370, and Asp537, disrupted homo-oligomerization of the Smad4/DPC4-C-terminal domain. Similar results were obtained with the full-length Smad4/DPC4 (Figure 3D). Conversely, the Drosophila /C. elegans developmental mutation Gly508Ser (Figure 2A) had no effect on homooligomerization (Figure 3D). This mutation maps to the L3 loop, which is the only portion of the loop/helix region not involved in the trimer interface.

If the Smad4/DPC4-C-terminal domain forms a trimer, then full-length Smad4/DPC4 should form a trimer as well. Figure 4A shows that recombinant full-length Smad4/DPC4, purified to near homogeneity, elutes from a gel-filtration column with an apparent

molecular size of ~180 kDa, consistent with the 181 kDa size calculated for the Smad4/DPC4 trimer. This large apparent size is likely the result of trimerization because the tumor-derived trimerinterface mutations reduce the apparent size by a factor of about three (Figure 4B). Conversely, the *Drosophila/C. elegans* developmental mutation Gly508Ser, which does not directly affect a trimer-interface residue, had no effect on the large apparent size of Smad4/DPC4 (Figure 4B). However, the Smad4/DPC4-C-terminal domain elutes as a monomer from a gel filtration column, consistent with residues N-terminal to the Smad4/DPC4-C-terminal domain contributing to homo-oligomerization.

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In principle, the full-length Smad4/DPC4 protein may assume an oligomeric state other than a trimer but still with a gel filtration mobility approximating that of a trimer. However, the *in vivo* and *in vitro* data with the trimer interface mutants, both with the C-terminal domain and the full-length proteins, strongly suggest that the trimeric protein-protein interface observed in the crystals is also the one that participates in homo-oligomerization *in vivo*.

Smad4/DPC4-C-terminal domain also supports hetero-oligomerization, shown by the co-immunoprecipitation overexpressed Smad4/DPC4-C-terminal domain and terminal domain from COS cells (Figure 3D), and by the association of Smad4/DPC4-C-terminal domain with Smad2-C-terminal domain in a native gel electrophoresis assay. Furthermore, the tumor-derived trimer interface mutations, as well as the developmental L3 loop mutation abolished hetero-oligomerization between the

Smad4/DPC4-C-terminal domain and the Smad2-C-terminal domain (Figure 3D). Similar results were obtained with the full-length The observation that the L3-loop developmental Smad4/DPC4. mutation, which did not significantly affect homo-oligomerization, disrupted hetero-oligomer formation, suggests that the L3 loop may hetero-oligomerization. participate in The observation that mutations that disrupted homo-oligomerization also disrupted hetero-oligomerization further suggest that homo-oligomer formation could be a prerequisite for hetero-oligomerization.

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Although several hetero-oligomerization models would be consistent with the available data, one model that is suitable, from a structural perspective, is the formation of a hetero-hexamer between Smad4/DPC4 and Smad2 trimers. As the trimer structure resembles a disk with the L3 loops forming undulations on the face of the disk (Figure 5A), this could allow two disks to come together face-to-face and interact via their L3 loops (Figure 5B), explaining why L3 loop mutations disrupt hetero-oligomerization. In this model, hetero-hexamer formation would also require homo-trimer formation, explaining how the tumorigenic mutations that disrupt homo-oligomerization can also disrupt the formation of the functional hetero-oligomeric complex and interfere with signal transduction.

#### EXAMPLE 7

Construction of expression vectors and Yeast two-hybrid system

To generate human Smad4 and Smad2 mutations, a fragment of the corresponding cDNAs was amplified by PCR. The

amplified region was subcloned into the full-length Smad4 or Smad2 in pCMV5 for mammalian cell transfection. The regions amplified by PCR and the presence of missence mutations were confirmed by sequencing.

LexA fusions were created in pBTM 116 and GAD fusions within pGAD424 (Clontech). Interactions were tested in the strain L40. Activation of the LexA operator-HIS3 reporter was assayed on media lacking histidine with increasing concentrations of 3-aminotriazole.

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#### EXAMPLE 8

Transfection, immunoprecipitation, immunoblot, and metabolic labeling

For Smad2/Smad4 homo- or hetero-complex analysis, COS cell were transiently transfected with the indicated constructs, and stimulated with 200 pM TGFβ1 for 1 hour. Cells were lysed in TNE buffer, immunoprecipitated with anti-Flag M2 monoclonal antibody (IBI; Eastman Kodak), and interacting proteins were detected by immunoblot with anti-HA monoclonal antibody 12CA5 (Boehringer Manheim) as described. Anti-Smad rabbit polyclonal antibody was raised against the full-length Smad1. To study interactions between N domain and C domain of Smad4 or Smad2, transiently transfected COS cells were lysed in LSLD buffer (50 mM Hepes, pH 7.4, 50 mM NaCl, 0.1% Tween 20, 10% glycerol, 1 mM DTT) containing protease and phosphatase inhibitors. Immuno-precipitation and immunoblot were done as described above. COS or R-1B/L17 cells transfected

with the indicated constructs were labeled with <sup>35</sup>S-methionine or <sup>32</sup>P-orthophosphate and visualized by electrophoresis and autoradiography.

5 <u>EXAMPLE 9</u>

## Functional assays

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For the animal cap assay, RNA (10 nl, 2 ng) was introduced in the animal pole of two-cell *Xenopus* embryos. Animal caps were explanted at blastula stage and cultured to tadpole stage. Total RNA from the harvested explants and control sibling embryos was extracted and RT-PCR was performed using muscle actin and EF-la primers. In the MDA-MB468 cell experiments, the amounts of transfected plasmids were adjusted in order to render the TGF\$\beta\$ response dependent on both Smad2 and Smad4. Luciferase and growth-inhibition assays were performed.

To investigate the domains involved in these interactions, various Smad4 fragments were tested as baits either against Smad4, to detect homo-oligomeric interactions, or against Smad2, to detect hetero-oligomeric interactions, in a yeast two-hybrid system. These experiments revealed that both the C domain and the domain/linker region can contribute to the homo-oligomeric interaction (Figure 6A). Full length Smad4 interacted with the N domain/linker region as a whole but not with these two regions expressed separately (Figure 6A). Full length interacted with its isolated C domain, albeit less strongly than with

itself (Figure 6A). Furthermore, isolated Smad4 C domain interacted strongly with itself (Figure 6A).

The higher affinity of full length Smad4 for itself than for its isolated C domain would result in the exclusion of the isolated C domain from full length homo-oligomeric complexes. Smad2 had a similar, albeit not identical pattern of homo-oligomeric interactions in yeast (Figure 6A). The homo-oligomeric interaction pattern of the Smads in yeast is consistent with a contribution of all three regions to the homo-oligomeric interaction, with the C domain providing the strongest interaction. Resolution of the crystal structure of the Smad4 C domain has revealed that this domain forms a homotrimer whose interfaces are the targets of cancer mutations.

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The Smad2-Smad4 interaction was detectable in yeast, and was particularly sensitive to deletions in the C domain (Figure 6A). Furthermore, Smad2 and Smad4 or their isolated C domains interacted strongly with each other's isolated C domain (Figure 6A). In contrast to its dependence on TGFβ stimulation in mammalian cells, the Smad2-Smad4 interaction occurred spontaneously in yeast. This might be due to Smad phosphorylation by protein kinases in yeast. These interactions were also analyzed in mammalian cells by transfection of Smad2 and Smad4 fragments tagged with the Flag or HA epitopes.

The isolated C domains of Smad2 and 4 each formed homo-oligomers in COS cells, as determined by co-immunoprecipitation of differently tagged constructs (Figure 6C). The Smad2-Smad4 interaction in COS cells requires TGFβ receptor

stimulation (Figure 6D) whereas the homo-oligomeric interactions do not (Figure 6C). The C domains of Smad2 and Smad4 were necessary and sufficient for this interaction (Figure 6D).

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Interestingly, deletion of the N domain in either Smad2 or Smad4 caused a constitutive association with the full length version of the other protein, and this association was further enhanced by TGFβ action (Figure 6D). The expression level of the Smad2 and Smad4 deletion products was similar to that of the full length proteins (Figure 6B), arguing that the constitutive association was not due to a higher expression of the N domain deletion products. Thus, the results in yeast and COS cells combined suggest that the Smad2-Smad4 interaction is mediated primarily by the C domains, requires the integrity of these domains, and is inhibited by the presence of the N domains (Figure 6E).

Consistent with this inhibitory function, expression of the Smad4 N domain (Figure 7A) or the Smad2 N domain (Figure 7B) inhibited the association between full length as well as the C domain forms of Smad2 and Smad4 in COS cells. This effect is specific since the overexpression of N domains does not inhibit homooligomerization of the C domains (Figure 7C) or the expression level of cotransfected C domains.

Since the formation of the Smad2-Smad4 complexes requires Smad2 phosphorylation, which occurs at C terminal serines, the relationship between this process and the inhibitory effect of the N domain was determined. The Smad2 and Smad4 C domains spontaneously associated with each other and their association was

further stimulated by TGFβ (Figure 8A). Interestingly, the Smad2 C domain was phosphorylated in response to TGFβ to an extent similar to that of the full length Smad2 (Figure 8B). Cotransfection of a kinase inactive, dominant negative TGFβ type I receptor [TβR-I(K232R)] abolished the TGFβ-stimulated association of Smad2 and Smad4 C domains but not their constitutive association (Figure 8C). Thus, the Smad2-Smad4 interaction is independently stimulated by removal of the N domains and by phosphorylation of the Smad2 C domain.

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The ability to analyze the inhibitory function of Smad N domains led to the investigation of Smad2 and Smad4 products containing N domain mutations identified in human cancers. Smad4 mutations have been identified in pancreas, colon, esophageal, breast, ovary, and head and neck cancers and Smad2 mutations in colon and head and neck cancers. Most of the missense mutations are in the C domains of Smad2 or 4. The three-dimensional structure of the Smad4 C domain predicts that some of these mutations destabilize the core structure whereas others disrupt the C domain homotrimer interface, and others disrupt a putative Smad4-Smad2 interface. However, missense mutations have also been identified in the N domains of Smad2 and Smad4. These include the Smad2 Arg133Cys mutation identified in a colon carcinoma and the Smad2 Arg100Thr mutation identified in a pancreatic carcinoma. Interestingly, Arg133 in Smad2 corresponds to Arg100 in Smad4, both of which are located in a highly conserved region of the Smads, suggesting a selection for mutations at this residue in cancer.

That these N domain mutations inactivate the signaling function of Smad2 and Smad4 was confirmed. In the Xenopus embryo, injection of Smad2 transcripts mimics the ability of activin to induce dorsal mesoderm in ectodermal explants. The Arg133Thr mutation eliminated the ability of Smad2 to induce muscle actin, a paraxial mesoderm marker, in this assay (Figure 9A). In the human carcinoma cell line MDA-MB468, which has a Smad4 homozygous deletion and is, therefore, insensitive to TGFB, Smad4 transfection restores TGFB sensitivity and this is enhanced by cotransfection of Smad2. As measured using the TGF\$\beta\$ reporter gene construct 3TP-luciferase, the Arg mutation in either Smad2, Smad4 or in both eliminated the ability of the cotransfected constructs to restore TGFB responsiveness in these cells (Figure 9B). Moreover, Smad4 overexpression inhibits MDA-MB468 cell proliferation, and this activity was also disrupted by the Arg100Cys mutation (Figure 9C).

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To investigate the basis for the inactivation effect of these N domain mutations, the mutant Smad2 and Smad4 were transfected into COS cells and their expression level and interactions were determined. Both mutants were similar to their wild type counterparts with regards to expression levels (Figure 10A) and their ability to form homo-oligomers (Figure 10A). However, both mutants failed to form Smad2-Smad4 complexes in response to TGFβ (Figure 10B). Since the Smad2-Smad4 interaction is primarily a function of the C domains and since the N domains repress this

interaction, the inhibitory function of the wild type and mutant N domains was further investigated.

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When expressed as separate polypeptides in COS cells, the N domains of Smad2 or Smad4 associate with the corresponding C domain (Figure 10B), providing evidence for a direct interaction between the N and C domains of a Smad protein. In yeast, the isolated N domains of Smads 2 and 4 interacted weakly with their respective C domains and not at all with the full length proteins (see Figure 6A). The latter could be due to an interference by an intramolecular N domain-C domain interaction in the full length protein. The interaction between the isolated N and C domains in COS cells is specific since it is not observed between the N domain of one Smad and the C domain of the other (Figure 10B). Importantly, the Smad2 and Smad4 mutant N domains interacted corresponding C domains 18 and 22-fold more strongly than did the wild type N domains (Figure 10B). Since these mutations do not increase the expression level of the N domains, this increase in binding is likely to result from an increased affinity of the mutant N domains for the C domains. Furthermore, the mutant N domains were more potent than the wild type N domains at inhibiting the Smad2-Smad4 hetero-oligomerization (Figure 10C).

To determine the effect of the Smad4 N domain on the Smad2-Smad4 signaling function, its effect on the ability of Smad2 and Smad4 to activate the 3TP-luciferase reporter was tested. In agreement with previous studies, overexpression of Smad2 and Smad4 or of their C domains activated this reporter (Figure 10D).

Cotransfection of the Smad2 or Smad4 N domains significantly inhibited this effect (Figure 10D). Furthermore, the mutant N domains were more potent than the wild type N domains as inhibitors of Smad2/Smad4-mediated response (Figure 10D). Thus, the ability of a Smad N domains to bind to the corresponding C domain correlates with inhibition of Smad2-Smad4 interaction and signaling function. The Arg133Thr and Arg100Cys mutations increase the inhibitory function of the Smad2 N domain and the Smad4 N domain with either mutation leading to inactivation of Smad2-Smad4 signaling function. Since Smad4 is a shared partner of other Smads besides Smad2, the Arg100Thr mutation would disrupt the signaling function of these other Smads as well.

In sum, the present invention demonstrated that the N domain in Smad proteins directly interacts with and represses the effector function of the C domain. Furthermore, certain Smad2 and Smad4 mutations found in human cancer inactivate these proteins by augmenting the inhibitory function of the N domain. Previously characterized tumor suppressor mutations, including missense mutations in the C domains of Smads 2 and 4, act by disrupting protein stability or effector function. The present findings reveal a mechanism of tumor suppressor inactivation involving instead a gain of autoinhibitory function. Antagonists of Smad autoinhibitory function might be useful in reversing the effects of this type of mutation.

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#### **EXAMPLE 10**

## Expression vectors

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Human Smad1, Smad2 and Smad4 mutations were made by a PCR-based strategy as described. All PCR-generated fragments were subcloned into wild type Smads in CMV promoter-based mammalian expression vectors pCMV5 or pCS2 and verified by sequencing.

The transfection. metabolic labeling and immunoprecipitation studies were performed as follows. For in vivo labeling with [35S]methionine or [32P]orthophosphate and for coimmunoprecipitation studies, cells were transiently transfected by the DEAE-dextran method as described above. To examine the phosphorylation of Flag-tagged Smad1 and Smad2 constructs, R-1B/L17 cells were co-transfected with either TBR-I or BMPR-IB and BMPR-II. Forty to 48 hours after transfection, cells were washed and preincubated with phosphate-free media for 1 hour. The cells were then incubated with the same phosphate-free media containing 1 mCi/ml [32P]phosphate for 2 hours at 37°C and then stimulated with either TGF-\(\beta\)1 (1 nM) or BMP4 (10 nM) for 30 minutes. Subsequently, labeled and ligand-stimulated cells were lysed in TNE buffer (10 mM Tris, pH 7.8; 150 mM NaCl; 1 mM EDTA; 1% NP40) containing protease and phosphatase inhibitors, and the lysates were subjected to immunoprecipitation with anti-Flag M2 monoclonal antibody (IBI; Eastman Kodak). Protein expression of Smads was determined either by metabolic labeling or western blotting. COS-1 cells that have been transiently transfected for 40-48 hours were washed and preincubated in methionine-free media and then labeled

with trans-[35] methionine for 3 hours. Lysis and immunoprecipitation were performed as for [35P] phosphate-labeled cells. Immunoprecipitates were visualized by SDS-polyacrylamide gel electrophoresis (SDS-PAGE) followed by autoradiography. For western blotting, a fraction of the total cell lysate was separated by SDS-PAGE and assayed by immunoblotting as indicated.

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For Smad4 association studies, Flag-tagged Smad1 or Smad2 constructs were transiently co-transfected with HA-tagged Smad4 into COS-1 cells. Forty to 48 hours after transfection, cells were washed in DMEM containing 0.2% fetal calf serum and treated with the indicated ligand (200 pM TGF-\beta1 or 5 nM BMP4). Following ligand-stimulation, cells were lysed in TNE buffer containing protease inhibitors. Cell lysates were then subjected to immunoprecipitation with anti-Flag M2 monoclonal antibody. Immunoprecipates were washed, separated by SDS-PAGE, and transferred to **PVDF** membranes (Immobilon-P; Millipore). HA-tagged Smad4 detected using anti-HA monoclonal antibody 12CA5 (Boehringer Mannheim), followed by donkey anti-mouse antibody conjugated with horseradish peroxidase (Sigma) and chemiluminescence (ECL, Amersham).

COS-1 cells transiently transfected for 40-48 hours by the DEAE-dextran method were affinity-labeled with [1251]TGF-β as described. Briefly, cells were preincubated at 37°C in Krebs Ringer Hepes (KRH) buffer containing 0.5% bovine serum albumin (BSA), washed with cold KRH/0.5% BSA, and affinity labeled using 200 pM [1251]TGF-β in KRH/0.5% bovine serum albumin (BSA) for 3.5 hrs at

Then, the cells were washed four times in ice-cold KRH 4°C. containing 0.5% BSA and once more with KRH alone. Subsequently, cell-surface bound [125]TGF-β was cross-linked to the receptor complex by incubation for 15 minutes at 4°C with 60 mg/ml disuccinimidyl suberate in KRH; cross-linking was terminated washing the cells twice with ice-cold STE (0.25 M sucrose, 10 mM Tris-HCl, pH 7.4 and 1 mM EDTA). Cells were then lysed in TNT [20] mM Tris-HCl, pH 7.4, 150 mM NaCl, 1% Triton X-100 (v/v)]<sup>37</sup> containing protease and phosphatase inhibitors and the cell lysate subjected to anti-Flag immunoprecipitation. Labeled receptor complexes in the immunoprecipitates and in the total cell lysates were then visualized by separation on SDS-PAGE and autoradiography.

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HepG2 cells were transfected overnight using the calcium phosphate-DNA precipitation method. Twenty-four hours after transfection, cells were transferred onto chamber slides (Nunc, Inc.). Forty to 48 hours post-transfection, cells were stimulated with 5 nM BMP4 or 1 nM TGF-β for 30 minutes and processed for immunofluorescence. Immunostaining was performed using anti-Flag M2 monoclonal antibody and FITC-conjugated secondary antibodies (Pierce).

The present invention shows that the L3 loop in the C domain of receptor-regulated Smads is crucial for their specific interaction with the TGF- $\beta$  and BMP receptors. Signal transduction specificity in the TGF- $\beta$  system was determined by ligand activation of a particular receptor complex which then recruits and

phosphorylates a subset of Smad proteins including Smads 1 and 2. These then associate with Smad4 and move into the nucleus where they regulate transcription. A discrete surface structure was identified in Smads 1 and 2 that mediates and specifies their receptor interactions. This structure is the L3 loop, a 17-amino acid region that, according to the crystal-structure of Smad4, protrudes from the core of the conserved Smad C-terminal domain. The L3 loop sequence is invariant among TGF-β-activated Smads (Smads 2 and 3) and BMP-activated Smads (Smads 1, 5, 9 and Mad) but differs at two positions between these two groups. Switching these two amino acids switches Smad1 and Smad2 activation by BMP and TGF-β, respectively. These studies identify the L3 loop as a critical determinant of specific Smad-receptor interactions.

#### 15 **EXAMPLE 11**

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## C-tail is dispensable for Smad2 association with the TGF-B receptor

Receptor-regulated Smads are phosphorylated activated receptors at conserved C-terminal serine According to the crystal structure of the Smad4 C-domain, thought to be conserved in the receptor-regulated Smads, these residues are located at the end of an 11-amino acid region (here referred to as the "C-tail") following  $\alpha$ -helix 5 (Figure 11A). As a substrate for the TGFβ type I receptor kinase, the C-tail might mediate the observed docking of Smad2 to the receptor complex. This possibility was examined by testing the receptor-binding activity of a Smad2 construct lacking the C-tail [Smad2(1-456)]. Receptor-binding

activity was assayed by co-transfection of TBR-I, TBR-II and Flag epitope-tagged Smad2 constructs into cells, then affinity-labeling the receptors by crosslinking to bound 125I-TGF-β1, and finally coimmunoprecipitating the labeled receptors with Smad2 via the Flag Surprisingly, the receptor interaction was epitope (Figure 12A). stronger with Smad2(1-456) than with wild type Smad2 (Figure 12A), indicating that removal of the C-tail increased the Smad2receptor interaction. This suggests that the physical contact between the C-tail of Smad2 and the catalytic cleft of the TBR-I kinase during the phosphotransfer reaction does not contribute significantly to Smad-receptor association. Smad2 docking to the receptor must therefore be mediated by a region of Smad2 other than the C-tail. The interaction between the TGF-\( \beta\) receptor complex and Smad2 is increased when TBR-I is made catalytically inactive by a mutation in the kinase domain or the C-terminal phosphorylation sites in Smad2 are eliminated by mutation to alanine [see Figure 12A, Smad2(3A) In light of the observation that removal of the C-tail increases the receptor interaction, these results suggest that docking is inhibited when the C-tail is phosphorylated.

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#### EXAMPLE 12

## The Smad2 Cdomain associates with the receptor complex

In order to localize the region of Smad2 required for association with the receptor, various Smad2 deletion mutants were tested for receptor binding activity (Figure 13). To facilitate the analysis without altering the C-terminus of Smad2, the kinase

defective TBR-I(KR) receptor construct was used, taking advantadge of its enhanced Smad2 binding phenotype. Deleting half of the N domain [Smad2(100-467) construct] or the entire N domain [Smad2(186-467)] had no appreciable effect on Smad2-receptor association. Consistent with this, the N domain (1-185) alone had no detectable affinity for the receptor complex. Furthermore, the C domain alone [Smad2(248-467)] was still capable of associating with the receptor complex, albeit more weakly. This could be due to the fact that the C domain forms homo-oligomers less stably than the full-length protein and that this homomeric complex might cooperatively associate with the receptor complex. As with the fulllength Smad2, the C domain interacted with the wild type TBR-I more stably when the C-terminal phosphorylation sites of Smad2 were mutated [Smad2(248-467/3A) construct] (Figure 13).

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### EXAMPLE 13

## L3 loop involvement in Smad2 docking

Given these results, this search for a critical determinant of receptor docking focused in the C domain of Smad2 excluding the C-tail. Two missense mutations in this region inhibit receptor-mediated phosphorylation. A colorectal tumor-derived mutant form of Smad2 with an aspartic acid to glutamic acid mutation (D450E) is defective in receptor-dependent phosphorylation (Figure 12B). However, this mutant was able to bind to the receptor as effectively as did the Smad2(3A) mutant (Figure 12A), suggesting that the

D450E mutation interferes with Smad2 phosphorylation and, as a result, enhances Smad2 binding to the receptor.

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A different result was obtained with another mutant, Smad2(G421S), a highly conserved glycine residue whose mutation to serine in Drosophila Mad or to aspartic acid in Caenorhabditis elegans Sma-2 causes null or severe developmental phenotypes. corresponding mutation in Smad1 inhibits BMP-induced phosphorylation of Smad1. In Smad2, the (G421S) mutation inhibited TGF-\u03b3-dependent phosphorylation (Figure 12B). Unlike the D450E mutation, however, the G421S mutation inhibited Smad2 binding to the receptor (Figure 12A). This suggested that Gly421 is involved, directly or indirectly, in Smad2 association with the receptor, and mutation of this residue may inhibit phosphorylation by preventing this association.

Gly421 is located in a highly conserved segment of the Smad2 C domain (Figure 11A). The crystal structure of Smad4 C domain reveals that this segment forms a solvent-exposed loop, the L3 loop, protruding from the β-sandwich core structure of the C domain (Figure 11B). The L3 loop is predicted to participate in Smad interaction with other proteins. To show that the intergrity of the L3 loop is required for Smad2-receptor association, various residues that are absolutely conserved in this loop (G423, Y426, and RQ428,429; see Figure 11A) were substituted with alanine. Gly423 of Smad2 corresponds to Gly348 in Sma-3, which is converted to Arg in a developmental mutant allele. As inferred from the Smad4 crystal structure, these mutations should not destabilize the folding

of Smad2. These mutants were indistinguishable from the wild type Smad2 in their expression levels and their ability to form homooligomers (TABLE II). However, these mutations diminished (G423A) or abolished (Y426A and RQ428, 429AA) Smad2 binding to the TGF-β receptor complex. Defective receptor binding was accompanied by defective TGF-β-induced phosphorylation and defective association with Smad4 as measured by co-immunoprecipitation with a cotransfected epitope-tagged Smad4 construct.

10 TABLE II

Properties of Smad2 L3 loop Mutants

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L3 loop Mutation	Expression Level	Homo- oligomer	Receptor Binding	TGFβ-induced Phosphorylation	Smad4 Binding		
Wild	+	+	+++	+	+++		
Type							
G421S	+	+	+/-	-	+/-		
G423A	+	+	+/-	n d	+/-		
A424P	+	+	+++	+	+++		
Y426A	+	+	-	-	-		
R427P	+	+	-	-	-		
R427A	+	+	-	n d	-		
RQ428,	+	+	-	-	-		
429AA							
T432K	+	+	-	-	•		
T432A	+	+	-	n d	+/-		
S433A	+	+	+/-	n d	+/-		

Table II: Properties of L3 loop mutants of Smad2. The

15 expression level of Flag-tagged Smad2 constructs was determined by
anti-Flag immunoblotting. Homo-oligomeric Smad2 interactions were

assessed by co-transfection of Flag-tagged and HA-tagged version of each construct. Smad4 binding to Smad2 was determined by cotransfection of Flag-tagged Smad2 constructs and HA-tagged Smad4. In both cases, cell lysates were immunoprecipitated with anti-Flag antibody and the precipitates immunoblotted using anti-HA antibody. Receptor binding was determined by the level of 125I-TGFβ1-labeled receptors that was co-immunoprecipitated with Flagtagged-Smad2 following two co-transfection schemes: defective TBR-I with full-length Smad2 constructs or wild type TBR-I with C-tail deletion versions of each Smad2 construct. The two schemes yielded similar results with each Smad2 transfection mutant. TGF-\$1-stimulated phosphorylation of Flag-tagged Smad2 constructs was determined. In the binding assays, +++ indicates a wild type level of binding, +/- indicates a binding level 5-fold less than wild type, and - indicates no detectable binding. nd, not determined.

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The effect of these mutations strongly suggested that the L3 loop plays a crucial role in mediating Smad2-receptor interactions. Several other mutations in the L3 loop also inhibited Smad2 association with the receptor. These include R427P, R427A, T432K, T432A and S433A (TABLE II). Various highly conserved residues in other regions of the Smad2 C domain that are surface-exposed as predicted from the tertiary structure of the the related Smad4 C domain were also mutated. Mutations in  $\alpha$ -helix 2 (P360R; QRY364-366YHH; W368F), in  $\alpha$ -helix 3 (A392Q), and in  $\alpha$ -helix 4

(A404T; Q407E) did not diminish the binding of Smad2 to the receptor complex, suggesting that the integrity of these other regions is not essential for Smad-receptor association.

# 5 EXAMPLE 14

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# The L3 loop specifies Smad-receptor interactions

A sequence comparison of the TGF-\(\beta\)-activated Smads (Smads 2 and 3) and the BMP/Dpp-activated Smads (Smads 1, 5, 9 and Mad) reveals that the L3 loop is invariant within each group but differs at two positions (corresponding to residues 427 and 430 in Smad2) between these two groups (Figures 11A and B). To determine whether the L3 loop can define the specificity of Smadreceptor interaction, the ability of Smadl and Smad2 to associate with the TGF- $\beta$  receptor complex was first compared (Figure 14A). The relative binding of Smad1 versus Smad2 to the TGF-B receptor complex was assessed in three different co-transfection schemes that optimize the TGF-\$\beta\$ receptor-Smad interaction: wild type Smad with kinase-defective receptor; wild type receptor with Smad C-tail deletion constructs; and, wild type receptor with Smad C-tail serine to alanine mutations. All three schemes yielded consistent results showing that Smad2 associated with the TGF-β receptor complex 5to15-fold more effectively than Smad1 (Figure 14A).

Whether the L3 loops of Smadl and Smad2 accounts for this differential affinity was tested. To this end, a Smad2 construct was created containing the Smadl L3 loop (by introducing the mutations R426H and T430D), and the reciprocal Smadl construct.

This Smad2 construct, hereafter referred to as Smad2(L1), had poor TGF-β receptor binding ability compared to Smad2, whereas the reciprocal construct Smad1(L2) was able to bind the TGF-β receptor complex as effectively as did Smad2 (Figure 14B). Switching the Ctails of Smads 1 and 2 in addition to the L3 loop [Smad1(LC2) and Smad2(LC1) constructs] had no additional effect on receptor binding (Figure 14B), consistent with the observation that the Smad2 C-tail does not contribute to docking to the receptor (Figure 12A). As expected, C-tail chimeras [Smad1(C2) and Smad2(C1) constructs] behaved like their wild type counterparts with regard to binding to the receptor. Thus, the Smad L3 loop critically determines the specificity of the Smad-receptor interactions.

## EXAMPLE 15

## 15 Switching Smad activation

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As shown in TABLE II, optimal receptor binding for Smad2 appeared to be necessary for the optimal phosphorylation of the C-tail (C). Consistent with this notion, TGF-β stimulation failed to phosphorylate Smad2(LC1) (Figure 15A), which is defective in binding to the TGF-β receptor, but phosphorylated Smad1(LC2) as effectively as it phosphorylated wild type Smad2 (Figure 15B). The requirements for Smad phosphorylation by activated BMP receptors was also investigated. Smad2(LC1) was phosphorylated in response to BMP receptor activation whereas Smad1(LC2) was not (Figure 15C). Thus, Smads 1 and 2 can be phosphorylated by a heterologous

receptor when they are allowed to dock to this receptor via a heterologous L3 loop.

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To determine whether optimal receptor binding is sufficient optimal C-tail phosphorylation, TGF-\beta receptormediated phosphorylation of the Smad1(L2) and Smad2(C1) constructs was examined. Both constructs bind to the TGF-\beta receptor but contain a Smad1 C-tail. Smad1(L2) was phosphorylated in response to TGF-β less extensively than were Smad2 or Smad1(LC2) (Figure 15B), even though all three constructs could bind to the TGFβ receptor equally well (see Figure 14B). On the other hand, Smad2(C1) was phosphorylated almost as efficiently as Smad2 in response to TGF-\(\beta\) (Figure 15A). Taken together, these data suggest that the non-conserved residues in the C-tail (see Figure 11A) have a limited influence on the phosphorylation of the C-terminal serines by the TGF-β receptor kinase.

To corroborate that the switch in receptor docking and phosphorylation specificity by introduction of the Smad2 L3 loop and C-tail into Smad1 resulted in the activation of Smad1(LC2) by TGF-β, the ability of this construct to associate with Smad4 was determined. Smad1(LC2) was able to associate with Smad4 in response to TGF-β whereas Smad2(LC1) was not (Figure 16A). Swapping the L3 loop and the C-tails between Smads 1 and 2 had no detectable effect on their ability to form homo-oligomers (Figure 16B). The ability of the TGF-β receptors and BMP receptors to induce Smad nuclear translocation was also switched in the Smad1(LC2) and Smad2(LC1) mutants (Figure 17). Like Smad2, Smad1(LC2) was translocated to

the nucleus in response to TGF-β but not BMP. On the other hand, like Smad1, Smad2 (LC1) was translocated to the nucleus in response to BMP but not TGF-β. Thus, the receptor input necessary to induce association of Smad1 or Smad2 with Smad4 and their movement to the nucleus is provided through a receptor interaction that is dependent on, and specified by, the L3 loop.

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Specificity is an essential property of signal transduction pathways. In the TGF-β signaling system, specificity is determined by ligand activation of a particular receptor combination which, in turn, recruits and phosphorylates a particular subset of Smad proteins. The present invention demonstrates the Smad-receptor interaction, the molecular basis for its specificity and identifies the L3 loop as a discrete surface structure in Smad proteins necessary for the Smad-receptor interaction and its specificity.

The differential ability of Smads 1 and 2 to associate with their known TGF-\(\beta\) receptor complex is consistent with responsiveness to these receptors: Smad2, which mediates TGF-B signaling, associates with the TGF-β receptor complex approximately 10-fold better than Smad1, which is primarily a mediator of BMP This receptor interaction is required for Smad2 signaling. phosphorylation since docking-defective mutants of Smad2 are not phosphorylated in response to TGF-β. However, the Smad2 phosphorylation sites themselves, along with the adjacent sequence in the 11-amino acid C-tail region, are dispensable for the receptor interaction. This conclusion is based on the observation that the TGFβ receptor associates with a Smad2 deletion mutant lacking the C-tail.

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These observations predict that a region other than the Ctail mediates Smad2 interaction with the activated TGF-B receptor complex. Since the isolated C domain of Smad2 still binds to the TGF-B receptor complex and, as with full-length Smad2, this interaction can be further enhanced by eliminating receptor-mediated Smad phosphorylation, a critical determinant of Smad docking resides in the C domain. Indeed, such a determinant was identified in a highly conserved region that, by analogy to the crystal structure of the Smad4 C domain, is predicted to form a highly solvent-exposed loop, the L3 loop, that is poised for protein-protein interactions. into the L3 loop, including Introduction of various mutations developmental mutations previously observed in Drosophila Mad and Caenorhabditis elegans Sma-2 and -3, diminishes the ability of Smad2 to associate with the TGF-\beta receptor complex. None of these mutations has appreciable effects on Smad2 expression level or its ability to homo-oligomerize, as predicted from the fact that the L3 loop is not part of the Smad C domain core structure.

The sequence of the L3 loop, which is invariant among TGF-β-activated Smads (Smads 2 and 3) and among Smads thought to be activated by BMP (Smads 1, 5, and 9) or Dpp (Mad), differs at two positions between these two groups. These two amino acids also differ in Smad4 as well as Smads 6 and 7 (Figure 11A). In Smad4, these two positions are highly exposed (Figure 11B), and the same is likely to occur in other Smads given their overall structural similarity to Smad4. As further testament to the importance of the L3 loop, switching these two amino acids in Smad1 and 2 induces a

gain or a loss, respectively, in their ability to bind to the TGF- $\beta$  receptor complex. This switch is reiterated in receptor-mediated phosphorylation of these Smads, indicating that the L3 loop-dependent receptor interaction is necessary and sufficient for receptor phosphorylation. The homologous C-tail containing the phosphorylation sites and adjacent sequence may ensure an optimal receptor-mediated phosphorylation. A switch in agonist-induced association with Smad4 and nuclear translocation accompanies this switch in phosphorylation.

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Unlike the receptor-regulated Smads, Smad4 lacks a Cterminal SS(V/M)S phosphorylation motif and does not appear to associate with the receptors on its own. What then is the function of the L3 loop in Smad4? Based on structural considerations and the observation that a mutation (G508S) in the Smad4 L3 loop abolishes the ability of Smad4 to associate with Smad2, the Smad4 L3 loop mediates the association with receptor-activated Smads. The importance of the Smad4 L3 loop for Smad2-Smad4 interaction has been shown by showing that mutations of other residues in the Smad4 L3 loop (Y513A; and RQ515,516AA) also lead to the loss of TGF-\u03b3-inducible Smad2-Smad4 association in transfected COS-1 cells. Smad4 is required for various responses to TGF-\$\beta\$, activin and BMP by acting as a partner for the corresponding receptor-activated Smads. In addition, Smad4 can associate with these Smads in yeast, suggesting that the interaction may be direct. Smad L3 loops, therefore, are implicated in two distinct types of interactions. Among the receptor-regulated Smads the L3 loop may mediate

Smad-receptor interactions, whereas the more divergent Smad4 L3 loop (see Figure 11A) may mediate Smad4 interaction with receptor-activated Smads. The L3 loop of receptor-regulated Smads may have a dual function as a receptor-interacting region and, upon phosphorylation of the C-tail, as a Smad4-interacting region.

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Since the C-tail of receptor-regulated Smads serves as a substrate for the type I receptor kinase, it must physically contact the receptor. But this interaction apparently does not contribute significantly to the stability of the interaction that precedes phosphorylation, at least as determined with Smad2 and the TGF-B receptor. In fact, the TGF-\beta receptor-Smad2 interaction is weakened upon phosphorylation by the receptor, as either phosphorylationdefective Smad2 mutants or a kinase-defective TGF-B type I receptor mutant enhances Smad-receptor association. It is not clear how Smad phosphorylation may promote its dissociation from the receptor. A gain of affinity for Smad4 might contribute to Smad2 dissociation from the receptor upon phosphorylation. However, the still showed an elevated receptor-binding Smad2(3A) mutant activity as compared to the wild type Smad2 in the Smad4-deficient colorectal carcinoma cell line SW480.7. Thus, an increased affinity for Smad4 may not be the only event driving dissociation of the phosphorylated Smad2 from the receptor complex.

Although two residues in the L3 loop are sufficient to dictate the specificity of the Smad-receptor interaction, the entire L3 loop may not be sufficient to fully support this interaction. It could be that a direct Smad-receptor interaction is weak and requires

oligomeric forms of both the receptors and the Smads for cooperative binding. Alternatively, the Smad-receptor interaction might be indirect, requiring a hitherto unidentified adaptor protein. Regardless of the mechanism, the evidence at hand identifies the L3 loop as a critical determinant of specific Smad-receptor interactions.

#### **EXAMPLE 16**

## Cell culture, Xenopus injections and animal cap assays

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R1B/L17 and COS-1 cells were maintained<sup>37</sup>. HepG2 cells were maintained in minimal essential medium (MEM, GIBCO-BRL) supplemented with 10% fetal bovine serum (FBS), nonessential amino acids and 2 mM sodium pyruvate. Mouse embryonal carcinoma P19 cells were cultured in DMEM medium supplemented with 10% FBS.

Receptor RNA (10 nl, 2 ng) was injected into the animal pole of two-cell embryos. Animal caps were explanted at the blastula stage and incubated to the tailbud stage (stage 28). RT-PCR of the indicated markers was performed.

#### EXAMPLE 17

## 20 Protein interaction, phosphorylation and immunofluorescence assays

Mutant receptor and Smad constructs were generated by PCR using appropriate oligonucleotides. Helix 2 exchange mutants

were generated by exchanging the six residues highlighted in the helix 2 region in Figure 20. Mutations were verified by DNA Wild-type and mutant receptors were C-terminally sequencing. tagged with a hemagglutinin (HA) epitope and were subcloned into the mammalian expression vector pCMV5. Cells were transiently transfected with the indicated constructs or empty vector by the DEAE-dextran method37. Phosphorylation of Smad1 and Smad2 was tested in R-1B/L17 cells by co-transfecting Flag-tagged Smad constructs and the indicated receptor constructs, labeling the cells with [32P]orthorphosphate for 2 h, followed by incubation with 1 nM TGF-B1 or 5 nM BMP2 for 30 min, and anti-Flag immunoprecipitation<sup>50</sup>. Expression levels of transfected proteins was determined by immunoprecipitation from [35S]methionine/cysteine labeled cells. Flag-tagged R-Smad interaction with HA-tagged Smad4 or myc-tagged Fast 1 was determined in COS-1 cells by anti-Flag immunoprecipitation and anti-HA or anti-myc western immunoblotting<sup>9,49</sup>. For Smad immunofluorescence assays, HepG2 cells were transfected overnight with DNA constructs as indicated, using the standard calcium phosphate-DNA precipitation method. Twenty-four h after transfection, cells were transferred chamber slides (Nunc, Inc.). Two days later, cells were stimulated with 5 nM BMP2 or 1 nM TGF-\beta1 for 1 h and processed for anti-Flag immunofluorescence<sup>50</sup>. The percentage of cells showing nuclear staining was determined by counting 200-300 positive cells.

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#### **EXAMPLE 18**

## Reporter Assays and Receptor Assays

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Activation of the p3TP-luciferase reporter construct<sup>32</sup> was analyzed in R1B/L17 cells<sup>37</sup>. To measure the activity of a Xvent2-luciferase reporter<sup>15</sup>, P19 cells were transfected with this construct, TβR-I and TβR-II. The next day, cells were incubated with 0.5 nM TGF-βI or 1 nM BMP2, and luciferase activity was measured 20 h later. To measure the activity of a Mix.2 ARE reporters (A3-CAT or A3-luciferase)<sup>45</sup>, R1B/L17 cells were transfected with these reporters, Fast1 and the indicated receptor constructs. The next day, cells were treated with 0.5 nM TGFβI or 1 nM BMP2 for 20 h and the reporter gene activity was determined<sup>49</sup>. A GAL4 DNA binding domain fusion with Fast1 was created by subcloning Fast1 into pGAD424 (Clontech). GAL4-Fast1 activation was determined in R-1B/L17 cells by cotransfection with the indicated constructs, and incubation with BMP2 for 14 h on the following day.

TGF-β1 and BMP2 were labeled with sodium [125I]<sup>67</sup>. To detect receptor-Smad interactions, COS-1 cells were transiently transfected with constructs that encode Smad1 and Smad2 lacking the last 11 amino acids [Smad1(1-454) and Smad2(1-456) constructs], and the indicated receptor constructs. After 40-48 h, cells were labeled by cross-linking to receptor-bound [125I]TGF-β1 or [125I]BMP2<sup>50</sup>.

#### **EXAMPLE 19**

# Determinants of specificity in the type I receptor

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The cytoplasmic domain of TGF-\beta family type I receptors was searched for regions that might determine the specificity of their interactions with R-Smads. One candidate was the GS domain, a 30 amino-acid region located just upstream of the kinase domain in all type I receptors<sup>63</sup>. The GS domain contains sites whose phosphorylation by the type II receptor activate the type I receptor kinase<sup>65</sup>. Phosphorylation sites in receptor tyrosine kinases function as docking sites for signal transduction molecules<sup>55</sup>. replacing the GS domain in the TGF-\$\beta\$ type I receptor, T\$\beta\$R-I, with the GS domain from one of the most divergent member of the TBR-I family in vertebrates, ALK2, did not alter the signaling specificity of TβR-I<sup>63</sup>. This result argued against a role of the GS domain in determining the specificity of receptor-Smad interactions.

A 9-amino acid segment in the receptor kinase domain, known as the "L45 loop", was also of interest (Figure 18A). It has been shown that replacement of all but the L45 loop in the kinase domain of TβR-I with the corresponding regions from ALK2 yields a construct that still mediates TGF-β responses<sup>38</sup>. As predicted from the conserved structure of protein kinases, the L45 loop links β-strands 4 and 5, and is not part of the catalytic center<sup>59</sup>. The L45 loop differs between type I receptors of different signaling specificity, such as the TGF-β receptors and the BMP receptors, but is

highly conserved between receptors of similar signaling specificity such as TβR-I and the activin receptor ActR-IB, or the BMP receptors from human (BMPR-IA and BMPR-IB) and *Drosophila* (Thick veins) (Figure 18A).

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To investigate the role of the L45 loop, TβR-I and BMPR-IB were used. The L45 loops of these two receptors differ by three non-conservative amino acid substitutions (Figure 18A). Constructs encoding these receptors with their L45 loops swapped were made by introducing N267I, D269G, N270T and T272S mutations in TβR-I, and the reciprocal mutations in BMPR-IB. These constructs showed a complete switch in their ability to activate Smadl and Smad2. Compared to the wild type receptors, TβR-I with the BMPR-I L45 loop [TβR-I(LB) construct] lost the ability to induce the formation of a Smad2-Smad4 complex and gained the ability to induce the formation of a Smad1-Smad4 complex (Figure 18B). The reciprocal pattern was observed with BMPR-IB containing the TβR-I L45 loop [BMPR-IB(LT) construct] (Figure 18B). These mutations also switched the ability of the receptors to induce translocation of Smadl and Smad2 into the nucleus (Figure 18C).

The L45 exchange mutations switched the signaling specificity of the receptors. BMPR-IB(LT) gained the ability to mediate TGF-β- and activin-like responses including activation of the 3TP-lux reporter construct, which contains a TGF-β response element from plasminogen activator inhibitor-1 and three AP-1 binding sites<sup>64</sup> (Figure 19A), and a reporter construct (A3-CAT) that contains

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activin- and TGF-\beta-responsive Fast1 binding sites from the Mix.2 promoter<sup>45</sup> (Figure 19B). TβR-I(LB) lost the ability to mediate these responses (Figure 19A and B) but gained the ability to mediate a BMP-like response, namely, activation of the Vent.2 promoter from Xenopus<sup>15</sup> in P19 mouse embryonal carcinoma cells (Figure 19C). Valine mutations of two conserved threonines (T272 and T274) at or near the TβR-I L45 loop did not impair 3TP-lux activation by TβR-I. Further evidence for a switch in signaling specificity was obtained using Xenopus embryo ectoderm explants. In these explants, TGF-B/activin signaling induces dorsal mesoderm and, indirectly, neural tissue via Smad216,40 whereas BMP signaling induces ventral mesoderm via Smad140.14.61. These effects can be observed using activated mutant forms of the corresponding type I receptors 41.58 However, an activated BMPR-IB receptor (see Figure 19D). containing the L45 loop from TBR-I [BMPR-IB(QD)(LT) construct] lost the ability to induce expression of the ventral mesoderm marker globin and gained the ability to induce the dorsal mesoderm marker muscle actin and the pan-neural marker NRP-1 (Figure 19D). The reciprocal construct, TβR-I(TD)(LB), showed an incomplete switch in signaling specificity in this assay system, losing the capacity to induce muscle actin without a gain of globin induction or a loss of NRP-1 induction (Figure 19D).

The switch in the signaling specificity of TβR-I(LB) and BMPR-IB(LT) correlated with a switch in their ability to recognize and phosphorylate Smads 1 and 2. The interaction between TGF-β

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family receptors and R-Smads is transient but can be visualized using mutant Smads lacking the receptor phosphorylation region<sup>50</sup>. As shown by co-precipitation of affinity-labeled receptors with Smads, TBR-I(LB) gained affinity for phosphorylation-defective Smadl and lost affinity for Smad2 compared to the wild-type receptors, whereas BMPR-IB(LT) lost affinity for Smad1 and gained affinity for Smad2 (Figure 20A). This switch extended to the pattern of receptor-dependent Smad phosphorylation. TBR-I and BMPR-I mediate C-terminal phosphorylation of Smad2<sup>30</sup> and Smad1<sup>48</sup>, respectively (see Figure 20B); basal phosphorylation (see Figure 20B) is due to MAP kinase action on inhibitory sites located in the central region of Smads<sup>47</sup>. In contrast to the effects of the wild type receptors, transfection of TBR-I(LB) elevated the phosphorylation of Smad1 whereas transfection of BMPR-IB(LT) elevated phosphorylation of Smad2 (Figure 20B). Interestingly, the increases in Smad phosphorylation caused by transfection of the L45 mutant rcceptors were ligand-independent. Indeed, TBR-I(LB) and BMPR-IB(LT) were hyperactive compared to the wild type receptors in in vitro kinase assays. The phenotype of a  $T\beta R-I$  allele containing a mutation (G261E) three residues upstream of the L45 loop had previously suggested that this region is involved in receptor activation<sup>62</sup>. However, despite their elevated kinase activity, the L45 mutant receptors had a clear switch in substrate specificity since TBR-I(LB) did not elevate Smad2 phosphorylation and BMPR-IB(LT) did not elevate Smad1 phosphorylation (Figure 20B). concluded that the subtype-specific residues in the receptor L45 loop

determine the specificity of Smad recognition, phosphorylation and activation.

#### EXAMPLE 20

## 5 Matching determinants of specificity in R-Smads

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The conserved C-terminal domain of R-Smad proteins, which is known as the "Mad homology-2" (MH2) domain, interacts with specific TGF-β family receptors and has specific effector functions. When expressed on its own in tissue culture cells or Xenopus embryos, the Smad2 MH2 domain is able to interact with the TGF-β receptor<sup>50</sup>, associate with Fast1<sup>49</sup> and generate TGF-β and activin-like effects<sup>16,42</sup>. These observations suggested that the receptor and DNA binding protein interactions of R-Smads are specified by determinants in the MH2 domain.

To search for such determinants, 21 amino acid residues of the MH2 domain that are not conserved between Smad1 and Smad2 but are highly conserved in Smads 1, 5, 8 and Mad, or in Smads 2 and 3 were investigated (Figure 21A). The location of these residues in the three-dimensional structure of the protein can be inferred from the crystal structure of the Smad4 MH2 domain<sup>57</sup>. The Smad4 MH2 monomer contains two β-sheets capped on one side by three α-helices (H3, H4 and H5) forming a bundle and, on the other side, by two large loops (L1 and L2) and an α-helix (H1). Smads

form homo-oligomers in the cell<sup>9,66</sup> and in solution<sup>57</sup>. In the crystal structure, the Smad4 MH2 domain forms a disc-shaped trimer, with the loop/helix region of one monomer forming an interface with the three-helix bundle of the next monomer (Figure 21B inset). Mutations in tumor-derived, inactive alleles of *Smad2* and *Smad4* often map to this interface<sup>57</sup>. At the amino acid sequence level, most of the structural elements of the Smad4 MH2 domain are conserved in the R-Smads (Figure 21A), which suggests that this three-dimensional structure is also conserved in R-Smads.

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10 Seven of the 21 subtype-specific amino acid residues (gray in Figure 21A) are clustered on the N-terminal side of the disc, near the point of connection to the N-terminal half of the Smad molecule; these residues are only partially exposed to solvent<sup>57</sup>. Two subtype-specific residues (yellow in Figure 21A) are located in αhelix 1, and six other (purple in Figure 21A) are at or near  $\alpha$ -helix 2, 15 which is highly exposed on the edge of the disc (Figure 21B). Of the remaining subtype-specific residues, two (red in Figure 21A) are located in the L3 loop, a structure protruding from each monomer on the C-terminal side of the disc (Figure 18B), and the last four (green 20 in Figure 21A) are located immediately upstream of the C-terminal receptor phosphorylation motif SS(V/M)S. Neither these four amino acids nor the phosphorylation motif itself are required for association with the TGF- $\beta$  receptor<sup>50,30</sup>.

Mutational analysis has shown that the L3 loop of Smad4 25 is essential for interaction with R-Smads<sup>57</sup> whereas the L3 loop of R-

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essential for Smads interaction with TGF-β receptors<sup>50</sup>. Furthermore, the two subtype-specific amino acids in this loop determine the specificity of the Smad-receptor interactions<sup>50</sup>. determine if the specificity of a R-Smad L3 loop matches specificity of the receptor L45 loop, it was investigated whether a Smad2 construct containing the Smad1 L3 loop sequence [Smad2(L1) construct] and the mutant  $T\beta R$ -I(LB) receptor construct would complement each other in the rescue of a TGF-\beta response. The association of Smad2 with Fast1 in response to agonist was used as a readout in these experiments. Formation of this complex recapitulates various additional signaling events (see Figure 18B). The Smad2(L1) construct bound Fast1 in response to BMP but not in response to TGF-β (Figure 22A), which is consistent with the ability of Smad2(L1) to recognize BMPR-IB but not T\u00e4R-I.50. T\u00e4R-I(LB) failed to mediate Smad2 association with Fast1. However, TBR-I(LB) mediated Smad2(L1) association with Fast1 (Figure 22B). Furthermore, the combination of TBR-I(LB) and Smad2(L1) rescued, partially at least, the ability to activate a Mix. 2 reporter construct in response to TGF- $\beta$  (Figure 22C). Therefore, the specificity of TGF- $\beta$ receptor-Smad interaction is determined by the L45 loop of the type I receptor and a complementary L3 loop in Smad2.

#### EXAMPLE 21

Determinants of Smad interaction with a DNA-binding partner

How a specific gene is targeted for activation by Smads has been delineated in the case of Mix.2. Activation of Mix.2 by activin or TGF-β requires the formation of a Smad2-Smad4-Fast1 complex which binds to a specific promoter sequence known as the "activin response element" (ARE)<sup>36,34,49</sup>. In this complex, the DNA binding domain of Fast1 mediates specific binding to the ARE<sup>36</sup> whereas the Smads act as transcriptional activators and enhancers of DNA binding<sup>49</sup>. The interaction between Smad2 and Fast1 is direct, as determined by their ability to interact as recombinant proteins in solution or in a yeast two-hybrid assays<sup>34</sup>.

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To identify a structural element that might specify the interaction of Smad2 with Fast1, it was investigated whether candidate Smad2 sequences introduced into Smad1 would allow it to recognize Fastl and activate a Mix.2 ARE reporter in response to BMP. The presence of six subtype-specific residues in the helix 2 of the MH2 domain (Figure 21A), and the prominent exposure of helix 2 on the edge of the MH2 trimer (Figure 21B) made this region a good candidate for this interaction. Exchanging the six subtype-specific helix 2 residues of Smad1 and Smad2 did not alter the specificity of their receptor interactions. Smadl containing the helix 2 sequence of Smad2 [Smad1(H2) construct] bound Smad4 in response to BMP, and the reciprocal construct, Smad2(H1), bound Smad4 in response to TGF-β (Figure 23A, upper panel). However, these helix 2 mutations switched the pattern of interactions with Fast1. Smad1(H2) gained the ability to associate with Fastl in response to BMP whereas Smad2(H1) failed to do so in response to TGF-\$\beta\$ (Figure 23A, lower

panel). Correlating with this switch, Smad1(H2) was able to mediate activation of a Mix.2 reporter in response to BMP whereas Smad2(H1) was unable to mediate activation of this reporter (Figure 23B). The Fast1 interaction specified by the Smad2 helix 2 was independent of the target promoter since Smad1(H2) was also able to activate a GAL4 reporter construct in cooperation with a Fast1-GAL4 DNA binding domain fusion (Figure 23C). These results suggest that  $\alpha$ -helix 2 of Smad2 is primarily responsible for the specificity for Fast1 and, as a result, the gene responses activated by the pathway. Extending these observations to the BMP pathway, Smad2(H1) gained the ability to mediate activation of a Vent.2 reporter in response to TGF- $\beta$  (Figure 23D).

#### EXAMPLE 22

### 15 Determinants of Specificity of TGF-β Signal Transduction

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Key determinants of specificity at three levels in the TGF-  $\beta$  and BMP signaling pathways have been identified. These determinants are encoded by specific amino acid residues in the L45 loop of the kinase domain in the type I receptors, and in the L3 loop and the  $\alpha$ -helix 2 of the MH2 domain in R-Smads. In each case, the residues involved are few and highly conserved in receptors or R-Smads that have similar signaling specificity. The interaction between these proteins may involve additional surface contacts, but results presented herein suggest that pathway specificity is largely

determined by these residues. Exchanging these residues at any of the three levels between TGF- $\beta$  and BMP pathway components switches the signaling specificity of these pathways.

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The L45 loop of type I receptor kinases had previously drawn attention because replacing the entire kinase domain except this loop in TBR-I with the corresponding regions from the functionally divergent receptor kinase ALK2 still allows mediation of TGF-B responses<sup>38</sup>. The L3 loop of Smads has drawn attention as a target of inactivating mutations in Drosophila and Caenorhabditis elegans Smad family members 18,19. As inferred from the effect of similar mutations in vertebrate Smads, the L3 loop participates in different interactions that are essential for signaling. In Smad4 the L3 loop is required for interaction with activated R-Smads<sup>57</sup>, whereas in R-Smads the L3 loop is required for interaction with the receptors and, furthermore, it specifies these interactions<sup>50</sup>. The present results show that matching combinations of L45 loops and L3 loops determine specificity of the receptor-Smad interaction. the Exchanging the subtype-specific residues in either the L45 loop or the L3 loop causes a switch in the specificity of this interaction, with an attendant switch in the signaling specificity of the pathway. As evidence of a functional match between a receptor L45 loop and a R-Smad L3 loop, the switch in the signaling specificity of a TGF-B receptor construct containing the BMP receptor L45 loop can be reversed by a Smad2 construct containing the matching L3 loop sequence from Smad1.

Results presented herein suggest that the interaction supported by the L45 and L3 loops achieves signal transduction by selectively increasing the affinity of a particular receptor kinase for a particular subtype of R-Smads. The docking interaction between receptors and R-Smads is independent of their catalytic interaction. The C-terminal SSXS phosphorylation motif of R-Smads and the the adjacent upstream sequence are neither required for association with the receptors in vivo nor for the specificity of this interaction<sup>50</sup>. However, effective R-Smad phosphorylation in vivo requires this docking interaction. Mutations that disrupt receptor docking strongly inhibit Smad phosphorylation and signal transduction. note, no stable interaction has been observed between the recombinant receptor kinase domains and Smads 1 or 2 in solution. Under these conditions, the TBR-I and BMPR-IB kinases phosphorylate both Smad1 and Smad2, and mutations in the L45 loop do not inhibit these reactions. The interaction supported by the L45 and L3 loops therefore might be cooperative, requiring the correct assembly of multivalent receptor complexes and R-Smad complexes in the cell.

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The present work also provides evidence that the choice of DNA binding partner and, consequently, the choice of target genes are determined by helix 2 in the MH2 domain of R-Smads. In the crystal structure of the Smad4 MH2 domain, helix 2 protrudes from the edge of the Smad trimer with several highly exposed residues. The sequence of helix 2 is divergent between R-Smads that mediate TGF-β (or activin) responses and those that mediate BMP responses,

but is highly conserved within each subgroup of R-Smads. Using as models the Mix.2 gene response to TGF- $\beta$  and the Vent.2 gene response to BMP, it was shown herein that the helix 2 of Smad2 and Smad1, respectively, determine the ability to mediate these responses. It was further shown that helix 2 from Smad2 specifies the selective interaction of Smads with the ARE-binding factor Fast1. Factors that mediates other Smad2- or Smad1-dependent gene responses remain to be identied. The ability of helix 2 to determine these interactions may provide ways to identify such factors. The role of helix 2 in Smad4 is also not known, although a mutation (R420H) in this region has been reported in lung carcinoma<sup>4</sup>.

The identification of determinants of specificity at three levels in TGF-β signal transduction suggests a general model for the organization of the selective protein-protein interactions that configure this signaling network (Figure 24). The determinants of specificity identified herein segregate the TGF-β and BMP pathways from each other. Still, each pathway can generate different responses in different cell types. Specificity at that level may depend on the repertoire of gene-targeting factors that the Smad complex encounters in the nucleus of a given cell.

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  - 64. Wrana, J.L., et al. 1992. Cell 71: 1003-1014.
  - 65. Wrana, J.L., et al. 1994. Nature 370: 341-347.
  - 66. Wu, R.-Y., et al. 1997. Mol. Cell. Biol. 17: 2521-2528.
  - 67. Cheifetz, S., et al. 1990. J. Biol. Chem. 265: 20533-20538.
- Any patents or publications mentioned in this specification are indicative of the levels of those skilled in the art to which the invention pertains. These patents and publications are herein incorporated by reference to the same extent as if each individual publication was specifically and individually indicated to be incorporated by reference.

One skilled in the art will readily appreciate that the present invention is well adapted to carry out the objects and obtain the ends and advantages mentioned, as well as those inherent therein. The present examples along with the methods, procedures, treatments, molecules, and specific compounds described herein are presently representative of preferred embodiments, are exemplary, and are not intended as limitations on the scope of the invention. Changes therein and other uses will occur to those skilled in the art which are encompassed within the spirit of the invention as defined by the scope of the claims.

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# SEQUENCE LISTING

	(1)	GENER	RAL IN	FORMATION:
		(i)	APPL	LICANT: Massagué, et al.
		(ii)	TITL	E OF INVENTION: Methods of Inhibiting or
5			Enhai	ncing the TGFβ-SMAD Signaling Pathway
		(iii)	NUM	BER OF SEQUENCES: 25
		(iv)	CORE	RESPONDENCE ADDRESS:
			(A)	ADDRESSEE: Benjamin Aaron Adler, Ph.D. J.D.
			(B)	STREET: 8011 Candle Lane
10			(C)	CITY: Houston
			(D)	STATE: Texas
			(E)	COUNTRY: United States of America
			(F)	ZIP: 77071
		(v)	COM	PUTER READABLE FORM:
15			(A)	MEDIUM TYPE: 1.44 Mb floppy disk
			(B)	COMPUTER: Apple Macintosh
			(C)	OPERATING SYSTEM: Macintosh
			(D)	SOFTWARE: Microsoft Word for Macintosh
		(vi)	CURI	RENT APPLICATION DATA:
20			(A)	APPLICATION NUMBER:
			(B)	FILING DATE:
			(C)	CLASSIFICATION:
		(vii)	PRIO	R APPLICATION DATE:
			(A)	APPLICATION NUMBER:
25			(B)	FILING DATE:
		(viii)	ATTO	DRNEY/AGENT INFORMATION:
			(A)	NAME: Benjamin Aaron Adler, Ph.D.
			(B)	REGISTRATION NUMBER: 35,423
			(C)	DECEDENCE/DOCKET NITIMBED, DOOLO

TELECOMMUNICATION INFORMATION:

(ix)

30

				(A)	TEI	_EPH	ONE	: (7	13)	777-2	2321				
				(B)	TEI	LEFA	X:	(713)	777	-690	8				
5	(2) INFORMATION FOR SEQ ID NO: 1														
	(i) SEQUENCE CHARACTERISTICS:														
	(A) LENGTH: 234 amino acids														
	(B) TYPE: amino acid														
	(C) STRANDEDNESS:														
10	(D) TOPOLOGY: linear														
			(	ii)	MOI	ECU	LET	YPE:							
	(A) DESCRIPTION: peptide														
	(iii) HYPOTHETICAL: no														
	(iv) ANTISENSE: no														
15	(v) FRAGMENT TYPE: internal														
			(	vi)	ORIO	GINA	L SO	URCE	Ξ:						
			(	vii)	IMM	EDLA	TE S	OUR	CE:						
			(	viii)	POS	OIT	N IN (	GENO	OME:						
			<b>(</b> i	ix)	FEA?	TURE	<b>:</b> :								
20			(:	x)	PUB!	LICA	MOIT	INF	ORM	(ATI	ON:				
			(:	xi)	SEQU	JENC	E DE	SCRI	РПО	N:	SE	Q ID	NO.:	1	
	Ala	Pro	Glu	Tyr	Trp	Cys	Ser	Ile	Ala	Tyr	Phe	Glu	Met	Asp	Val
	Cln	17a ]	C1	<b>01.</b> .	5 m	DI			_	10					15
25	GIII	vai	GIY	GIU	Thr 20	Pne	Lys	Val	Pro	Ser 25	Ser	Cys	Pro	Ile	
	Thr	Val	Asp	Gly	Tyr	Val	Asp	Pro	Ser		Glv	Asp	Ara	Phe	30 Cvs
					35					40			9		45
	Leu	Gly	Gln	Leu	Ser	Asn	Val	His	Arg		Glu	Ala	Ile	Glu	Arg
					50					55					60

	Ala	Arg	Leu	His		Gly	Lys	Gly	Val		Leu	Glu	Cys	Lys	_
					65					70			_		75
	Glu	Gly	Asp	Val	_	Val	Arg	Cys	Leu		Asp	His	Ala	Val	
_	_		_	_	80	_				85		_			90
5	Val	Gln	Ser	Tyr	_	Leu	Asp	Arg	GIu		GIY	Arg	Ala	Pro	
					95 -		_	_	_	100	_		_		105
	Asp	Ala	Val	His	_	Ile	Tyr	Pro	ser		lyr	iie	rys	Val	
	_	•		G1	110	*** =		Q3	<b>3</b> 4 - 4-	115	Q1	<b>0</b> 1		<b>.</b>	120
1.0	Asp	Leu	Arg	GIN		His	Arg	GIN	Met		GIN	GIN	Ala	Ala	
10	. 7 -	01×	77.	71-	125	77.	77-	Cl.	71-	130	* 1 ~	777	71-	C1	135
	Ala	GIII	Ala	Ala	140	Ala	Ата	GIII	Ala	145	Ата	vaı	АТА	GIY	150
	Tla	Pro	Glv	Pro		Ser	Val	Glv	Glv		Δla	Pro	Δla	Tle	
	116	110	CLY	110	155	DCI	V 4.1	011	013	160		110		110	165
15	Leu	Ser	Ala	Ala		Gly	Ile	Glv	Val		Asp	Leu	Ara	Ara	
10					170	3				175			3		180
	Cys	Ile	Leu	Arg	Met	Ser	Phe	Val	Lys	Gly	Trp	Gly	Pro	Asp	Tyr
	-				185					190					195
	Pro	Arg	Gln	Ser	Ile	Lys	Glu	Thr	Pro	Cys	Trp	Ile	Glu	Ile	His
20					200					205					210
	Leu	His	Arg	Ala	Leu	Gln	Leu	Leu	Asp	Glu	Val	Leu	His	Thr	Met
					215					220					225
	Pro	Ile	Ala	Asp	Pro	Gln	Pro	Leu	Asp						
					230										
25		** **							_						
	(3)	INF	)RML	ATIO	N FC	R SE	QШ	NO:	2						
	(i) SEQUENCE CHARACTERISTICS:														
					(A)	LE	<b>VGTI</b>	H: 19	97 ar	nino	acids				
					(B)	TY	PE:	amin	o acid	d					
30					(C)	STF	RAND	EDN	ESS:						
					(D)	TO	POLO	OGY:	linea	ır					
			G	ii)		ECUI									
			١,	/											

(A) DESCRIPTION: peptide

			(	111)	HYF	OTH	ETIC	AL:	no						
			(	iv)	AN	TISEN	NSE:	no							
			(	v)	FRA	GME	ENT T	ГҮРЕ	: in	terna)	l				
			(	vi)	ORI	GINA	L SO	URCI	E:						
5			(	vii)	IMM	ŒDL/	ATE S	OUR	CE:						
			(	viii)	POS	ITIOI	V IN	GEN	OME:						
				ix)		TURE									
				x)				N INTE	FORM	<i>የ ለ</i> ጥፕ	ONI.				
			•									. O. TD		_	
10	Larg	uic	`	xi)					PTIO			Q ID			
10	nys	nrs	ΙΙĐ	Cys	5	TIE	vaı	Tyr	туг	10	Leu	Asn	Asn	Arg	Val
	Gly	Glu	Ala	Phe		Ala	Ser	Ser	Thr		Val	Leu	Val	Asp	15 Gly
					20					25					30
	Phe	Thr	Asp	Pro	Ser	Asn	Asn	Lys	Asn	Arg	Phe	Cys	Leu	Gly	Leu
15					35					40					45
	Leu	Ser	Asn	Val		Arg	Asn	Ser	Thr		Glu	Asn	Thr	Arg	Arg
	714 -	<b>T</b> 1_	01	T	50	** 7		_	_	55 _					60
	птъ	TIE	Gly	ьуs	65	vaı	HIS	Leu	Tyr	7yr 70	Val	Gly	Gly	Glu	
20	Tyr	Ala	Glu	Cys		Ser	Asp	Ser	Ser		Phe	Val	Gln	Ser	75 Ara
				_	80		-			85		• • • •	0111	DCI	90
	Asn	Cys	Asn	Tyr	His	His	Gly	Phe	His	Pro	Thr	Thr	Val	Cys	Lys
					95					100					105
25	Ile	Pro	Ser	Gly		Ser	Leu	Lys	Ile		Asn	Asn	Gln	Glu	Phe
25	71 a	C15	T 011	Ton	110	C1	0	*** 7		115	~3				120
	AId	GIII	Leu	ьец	125	GIII	Ser	vai	Asn	130	GIY	Phe	GIu	Thr	
	Tyr	Glu	Leu	Thr		Met	Cys	Thr	Ile		Met	Ser	Phe	Val	135
					140		-			145				, u _	150
30	Gly	Trp	Gly	Ala	Glu	Tyr	His	Arg	Gln	Asp	Val	Thr	Ser	Thr	
					155					160					165
	Суѕ	Trp	Ile	Glu		His	Leu	His	Gly	Pro	Leu	Gln	Trp	Leu	Asp
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5	(4) INFORMATION FOR SEQ ID NO: 3															
			(	i)	SEQ	SEQUENCE CHARACTERISTICS:										
					(A)	A) LENGTH: 196 amino acids										
					(B)	(B) TYPE: amino acid										
					(C) STRANDEDNESS:											
10					(D) TOPOLOGY: linear											
	(ii) MOLECULE TYPE:															
	(A) DESCRIPTION: peptide															
	(iii) HYPOTHETICAL: no															
			(:	iv)	ANTISENSE: no											
15			(	v)	FRAGMENT TYPE: internal											
			(	vi)	ORIGINAL SOURCE:											
			(	vii)	IMMEDIATE SOURCE:											
			(	viii)	POSI	TION	IN C	GENC	ME:							
			(i	ix)	FEA7	TURE	<b>:</b>									
20			(:	x)	PUB	LICA	TION	INF	ORM	ATIO	ON:					
			(:	xi)	SEQU	JENC	E DE	SCRI	PTIO	N:	SE	QID	NO.:	3		
	Ala	Phe	Trp	Cys		Ile	Ala	Tyr	Tyr	Glu	Leu	Asn	Gln	Arg	Val	
					5					10					15	
	Gly	Glu	Thr	Phe		Ala	Ser	Gln	Pro	Ser	Leu	Thr	Val	Asp	Gly	
25					20					25					30	
	Phe	Thr	Asp	Pro		Asn	Ser	Glu	Arg	Phe	Cys	Leu	Gly	Leu	Leu	
					35					40					45	
	Ser	Asn	Val	Asn	Arg	Asn	Ala	Thr	Val	Glu	Met	Thr	Ara	Ara	His	

Ile Gly Arg Gly Val Arg Leu Tyr Tyr Ile Gly Gly Glu Val Phe

	Ala	Glu	Cys	Leu	Ser 80	Asp	Ser	Ala	Ile	Phe 85	Val	Gln	Ser	Pro	
	Cvs	Asn	Gln	Ara	Tyr	Glv	תיים	His	Pro		ጥከኍ	Val	C110	****	90
	<b>-</b> 21 -2			5	95	3		1120		100	****	Vai	Cys	БУS	105
5	Pro	Pro	Gly	Cys	Asn	Leu	Lys	Ile	Phe		Asn	Gln	Glu	Phe	
					110					115					120
	Ala	Leu	Leu	Ala	Gln	Ser	Val	Asn	Gln	Gly	Phe	Glu	Ala	Val	Tyr
					125					130					135
	Gln	Leu	Thr	Arg	Met	Cys	Thr	Ile	Arg	Met	Ser	Phe	Val	Lys	Gly
10	_				140			_		145					150
	Trp	GLy	Ala	Glu	Tyr	Arg	Arg	Gln	Thr		Thr	Ser	Thr	Pro	
	Ф~~	Tlo	Glu	Lou	155	T 011	ui.a	C111	Dwa	160	Q1	<b>m</b>	<b>.</b>	_	165
	TIP	116	GIU	ьеu	His 170	Leu	nis	GIY	Pro	175	GIN	Trp	ьeu	Asp	
15	Val	Leu	Thr	Gln	Met	Gly	Ser	Pro	Ser		Ara	Cvs	Ser	Ser	180
					185	-				190	5	0,0	501	501	195
	Ser														
	(F)	INTE	א א מר	י איז	NI EO	D CE	0 ID	NO	4						
20	(3)	IIALC			N FO		_								
20			(i	)	SEQL										
					(A)	LEN	IGTE	I: 19	96 an	nino	acids				
					(B)	TYF	E: a	amino	acid	l					
					(C)	STR	AND)	EDNE	SS:						
					(D)	TOF	OLO	GY:	linea	r					
25			(i:	i) :	MOLI	ECUL	ETY	PE:							
					(A)	DES	CRIE	TIO	N: p	eptid	e				
			(i	ii)	HYPO	OTHE	TICA	L: n	0						
			(i	v) .	ANTI	SENS	SE: n	10							
			(v	·) ]	FRAC	MEN	T T	YPE:	inte	rnal					
30			(v	i) (	ORIG	INAL	SOU	RCE:							
			(v	ii) l	IMME	EDIA.	ΓESC	URC	E:						
			(v	iii) l	POSIT	ΓΙΟΝ	IN G	ENO	ME:						

## (ix) FEATURE:

# (x) PUBLICATION INFORMATION:

			(:	xi)	SEQ	UENC	E DE	SCRI	PTIO	N:	SE	Q ID	NO.:	4	
	Ala	Phe	Trp	Cys	Ser	Ile	Ser	Tyr	Tyr	Glu	Leu	Asn	Gln	Arg	Val
5					5					10					15
	Gly	Glu	Thr	Phe	His	Ala	Ser	Gln	Pro	Ser	Met	Thr	Val	Asp	Gly
					20					25					30
	Phe	Thr	Asp	Pro	Ser	Asn	Ser	Glu	Arg	Phe	Cys	Leu	Gly	Leu	Leu
					35					40					45
10	Ser	Asn	Val	Asn	Arg	Asn	Ala	Ala	Val	Glu	Leu	Thr	Arg	Arg	His
					50					55					60
	Ile	Gly	Arg	Gly		Arg	Leu	Tyr	Tyr	Ile	Gly	Gly	Glu	Val	Phe
					65					70					<b>7</b> 5
1.5	Ala	Glu	Cys	Leu		Asp	Ser	Ala	Ile	Phe	Val	Gln	Ser	Pro	Asn
15	_				80					85					90
	Cys	Asn	Gln	Arg		Gly	Trp	His	Pro	Ala	Thr	Val	Cys	Lys	Ile
	_	_			95					100					105
	Pro	Pro	GIY	Cys		Leu	Lys	Ile	Phe	Asn	Asn	Gln	Glu	Phe	Ala
20	~ 7	_	_		110		_			115					120
20	Ala	Leu	Leu	Ala		Ser	Val	Asn	Gln	Gly	Phe	Glu	Ala	Val	
	C1	T	mla a a	2	125	<b>~</b>	em)			130					135
	GIII	Leu	Thr	Arg		Cys	unr	TTE	Arg	Met	Ser	Phe	Val	Lys	_
	Trn	Clar	ח [ ת	C1.,	140	7 ~~	7	<b>01</b> -	<b>m</b> \	145	<b>6</b> 13	_	_,	_	150
25	пр	GTĀ	ALA	GIU	155	AIG	Arg	GIN	Thr	Val	Tnr	Ser	Thr	Pro	_
23	Trn	T10	Glu	T.011		T.OU	Wic	Clu	Dro	160	C1-	<b></b>	T	•	165
	115	110	GIU	Dea	170	Бец	1112	GTĀ	PIO	Leu 175	GIII	Trp	Leu		
	Val	Leu	Thr	Gln		Glv	Ser	Pro	So~	Ile	7 22	C	Com		180
				<b></b>	185	7-7	501	-10	DET	190	AT 9	CYS	SeT.		
30	Ser									170					195

# (6) INFORMATION FOR SEQ ID NO: 5

## (i) SEQUENCE CHARACTERISTICS:

(A) LENGTH: 198 amino acids

			(B) TY	PE: ami	no ac	id					
			(C) ST	RANDEDI	VESS:						
			(D) TO	POLOGY	: line	ar					
		(ii)	MOLECU	LE TYPE:							
5			(A) DE	SCRIPTION	ON:	peptid	le				
		(iii)		ETICAL:		- •					
		(iv)	ANTISEN	NSE: no							
		(v)	FRAGME	ENT TYPE	E: inte	ernal					
		(vi)	ORIGINA	L SOURCI	Ξ:						
10		(vii)	IMMEDIA								
		(viii)	POSITION								
		(ix)	FEATURE								
		(x)	PUBLICA		ORM	ATIO	N·				
		(xi)	SEQUENC					O ID	NO.:	5	
15	Pro Lys	His Trp	Cys Ser								Ara
			5		_	10					15
	Val Gly	Glu Ala	Phe His	Ala Ser	Ser	Thr	Ser	Val	Leu	Val	Asp
	Clar Dho	Пb 3	20		_	25					30
20	GIY Phe	THE ASP	Pro Ser	Asn Asn	Lys	Ser .	Arg	Phe	Cys	Leu	
	Leu Leu	Ser Asn	Val Asn	Arg Asn	Ser		Ile	Glu	Asn	Thr	45 Ara
			50			55					60
	Arg His	Ile Gly	Lys Gly	Val His	Leu	Tyr '	Tyr	Val	Gly	Gly	Glu
25	Val m	71- Ol.	65 Com T	2		70		_			75
23	val Tyr	Ala Glu	Cys Leu 80	Ser Asp		Ser :	Ile	Phe	Val	Gln	
	Arg Asn	Cys Asn	Phe His	His Gly			Ser	Thr	Ser	Val	90 Cvs
			95	-		100			501	vui	105
	Lys Ile	Pro Ser	Ser Cys	Ser Leu	Lys	Ile I	Phe	Asn	Asn	Gln	Glu
30	<b>n</b> ) - "		110	_		115					120
	Phe Ala	GIn Leu	Leu Ala	Gln Ser			His	Gly	Phe	Glu	
			125			130					135

WO 99/01765 PCT/US98/13721 Val Tyr Glu Leu Thr Lys Met Cys Thr Ile Arg Met Ser Phe Val 140 145 150 Lys Gly Trp Gly Ala Glu Tyr His Arg Gln Asp Val Thr Ser Thr 155 160 Pro Cys Trp Ile Glu Ile His Leu His Gly Pro Leu Gln Trp Leu 5 170 175 180 Asp Lys Val Leu Thr Gln Met Gly Ser Pro Leu Asn Pro Ile Ser 185 190 195 Ser Val Ser (7) INFORMATION FOR SEQ ID NO: 6

- **SEQUENCE CHARACTERISTICS:** (i)
  - LENGTH: 197 amino acids (A)
- 15 (B) TYPE: amino acid
  - (C) STRANDEDNESS:
  - (D) TOPOLOGY: linear
  - MOLECULE TYPE: (ii)
    - (A) DESCRIPTION: peptide
- 20 (iii) HYPOTHETICAL: no
  - (iv) ANTISENSE: no
  - FRAGMENT TYPE: internal (v)
  - (vi) ORIGINAL SOURCE:
  - (vii) IMMEDIATE SOURCE:
- 25 (viii) POSITION IN GENOME:
  - (ix) FEATURE:
  - (x)PUBLICATION INFORMATION:
- (xi) SEQUENCE DESCRIPTION: SEQ ID NO.: 6 Ala Phe Trp Ala Ser Ile Ala Tyr Tyr Glu Leu Asn Cys Arg Val 30 10 15 Gly Glu Val Phe His Cys Asn Asn Asn Ser Val Leu Val Asp Gly 20 25 30

	Phe	Thr	Asn	Pro	Ser	Asn	Asn	Ser	Asp	Arg	Cys	Cys	Leu	Gly	Gln
					35					40					45
	Leu	Ser	Asn	Val	Asn	Arg	Asn	Ser	Thr	Ile	Glu	Asn	Thr	Arg	Arg
					50					55					60
5	His	Ile	Gly	Lys	Gly	Val	His	Leu	Tyr	Tyr	Val	Thr	Gly	Glu	Val
					65					70					75
	Tyr	Ala	Glu	Cys	Leu	Ser	Asp	Ser	Ala	Ile	Phe	Val	Gln	Ser	Arg
					80					85					90
1.0	Asn	Cys	Asn	Tyr		His	Gly	Phe	His	Pro	Ser	Thr	Val	Cys	Lys
10	_				95					100					105
	Ile	Pro	Pro	Gly		Ser	Leu	Lys	Ile	Phe	Asn	Asn	Gln	Glu	Phe
	_ •				110					115					120
	Ala	Gln	Leu	Leu		Gln	Ser	Val	Asn	Asn	Gly	Phe	Glu	Ala	Val
1.5	_		_		125					130					135
15	Tyr	Glu	Leu	Thr		Met	Cys	Thr	Ile	Arg	Met	Ser	Phe	Val	Lys
	01	_	<b>~</b> 1		140					145					150
	GIY	Trp	Gly	Ala		Tyr	His	Arg	Gln		Val	Thr	Ser	Thr	Pro
	0	<b></b>	~ 1	~1	155					160					165
20	Cys	Trp	Ile	GIU		His	Leu	His	Gly		Leu	Gln	Trp	Leu	Asp
20	T	**- 7	_	-1	170					175					180
	ьуs	val	Leu	Thr		Met	Gly	Ser	Pro		Asn	Ala	Ile	Ser	Ser
	t/a l	Cox			185					190					195
	Val	ser.													

# 25 (8) INFORMATION FOR SEQ ID NO: 7

- (i) SEQUENCE CHARACTERISTICS:
  - (A) LENGTH: 199 amino acids
  - (B) TYPE: amino acid
  - (C) STRANDEDNESS:
- 30 (D) TOPOLOGY: linear
  - (ii) MOLECULE TYPE:
    - (A) DESCRIPTION: peptide
  - (iii) HYPOTHETICAL: no

		(i	v)	ANT	ISEN	SE:	no									
		()	v)	FRA	GME	NT T	YPE	: inte	ernal							
		()	vi)	ORIC	INAI	L SOT	JRCE	i:								
		()	vii)	IMM	EDIA	TE S	OUR	CE:								
		(1	viii)	POSI	TION	I IN C	BENC	ME:								
		i.														
		•	·				INF	∩RM	ΑΤΙ	)N·						
											O ID	NO.	7			
Gln	Dhe	`	,	_							-			۲ <i>۲</i>		
GIII	rne	ILD	AIG	5	vai	per	IYL	17.1		Deu	ASII	1111	Arg	15		
Gly	Glu	Gln	Val	Lys	Val	Ser	Ser	Thr		Ile	Thr	Ile	Asp			
				20					25				_	30		
Phe	Thr	Asp	Pro	Cys	Ile	Asn	Gly	Ser	Lys	Ile	Ser	Leu	Gly	Leu		
				35					40					45		
Phe	Ser	Asn	Val		Arg	Asn	Ala	Thr		Glu	Asn	Thr	Arg	_		
•										_				60		
Hıs	Ile	GIY	Asn		Val	Lys	Leu	Thr		Val	Arg	Ser	Asn			
Ser	T.All	Dhe	λla		Care	Glu	Ser	) en	-	בות	Tlo	Dho	1727	75		
Ser	Dea	THE	ΛIα		Суз	Gra	Ser	dan		AIa	116	FIIE	vaı	90		
Ser	Ser	Asn	Cys		Tyr	Ile	Asn	Gly		His	Ser	Thr	Thr			
				95				_	100					105		
Val	Lys	Ile	Ala	Asn	Lys	Cys	Ser	Leu	Lys	Ile	Phe	Asp	Met	Glu		
				110					115					120		
Ile	Phe	Arg	Gln		Leu	Glu	Asp	Cys	Ser	Arg	Arg	Gly	Phe	Asp		
									130	_				135		
Ala	Ser	Phe	Asp		Gln	Lys	Met	Thr		Ile	Arg	Met	Ser			
170 l	T	C1**	Пххх		7 1 n	C1.,	Mb ess	C1-		01-	7	T7_ 7	m1	150		
vai	Lys	GIY	Trp		Ala	GIU	Tyr	GIII		GIN	Asp	vaı	Thr	165		
Thr	Pro	Cvs	Tro		Glu	Ile	His	Leu		Ala	Pro	Leu	Ala			
		-1-	12	170					175					180		
	Gly Phe Phe His Ser Val Ile Ala Val	Gly Glu Phe Thr Phe Ser His Ile Ser Leu Ser Ser Val Lys Ile Phe Ala Ser Val Lys	Gln Phe Trp Gly Glu Gln Phe Thr Asp Phe Ser Asn His Ile Gly Ser Leu Phe Ser Ser Asn Val Lys Ile Ile Phe Arg Ala Ser Phe Val Lys Gly	(viii) (ix) (x) (xi) Gln Phe Trp Ala Gly Glu Gln Val Phe Thr Asp Pro Phe Ser Asn Val His Ile Gly Asn Ser Leu Phe Ala Ser Ser Asn Cys Val Lys Ile Ala Ile Phe Arg Gln Ala Ser Phe Asp Val Lys Gly Trp	(vi)       FRAME         (vii)       ORIO         (viii)       IMM         (viii)       POSI         (ix)       FEAT         (xi)       SEQU         (xi)       SEQU         Gln       Phe       Ana       Thr         5       Gly       Ana       Thr         6       Ana       Val       Ana         Phe       Ana       Ana       Gly         Bhis       Ile       Ana       Ile         Bhis       Ile       Ana       Ile         Bhis       Ile       Ana       Ile         Bhis       Ile       Ana       Ile         Bhis       Ile       Ana       Ile	(v)       FRAGME         (vi)       ORIGINAL         (vii)       IMMEDIA         (viii)       POSITION         (ix)       FEATURE         (x)       PUBLICA         (xi)       SEQUENCE         Gln       Phe       Trp       Ala       Thr       Val         Gly       Glu       Gln       Val       Lys       Val         Gly       Glu       Glu       Val       Lys       Val         Phe       Asn       Val       Asn       Arg         Phe       Asn       Cys       Asn       Tyr         Ser       Leu       Phe       Ala       Glu       Cys         Ser       Asn       Cys       Asn       Lys         Val       Lys       Ile       Ala       Asn       Lys         Ile       Phe       Arg       Glu       Leu       Leu         Ile       Phe       Asp       Leu       Glu       Ala         Ile       Phe       Asp       Leu       Glu       Ala         Ile       Phe       Asp       Leu       Glu       Ala       Ala       Ala       Ala	(v)       FRAGMENT TO COLSTICAL SOLD         (vii)       IMMEDIATE SOLD         (viii)       POSITION IN COLSTION IN COLSTICATION (xi)         (x)       PUBLICATION (xi)         (xi)       SEQUENCE DE COLSTION (xi)         (xi)       (xi)       (xi)       (xi)       (xi)       (xi)       (xi)       (xi)       (xi)       (xi)       (xi) <th c<="" td=""><td>  (vi</td><td>  (v)   FRAGMENT TYPE: into</td><td> </td><td>(v) FRAGMENT TYPE: internal         (vi) ORIGINAL SOURCE:         (vii) IMMEDIATE SOURCE:         (viii) POSITION IN GENOME:         (ix) FEATURE:         (xi) SEQUENCE DESCRIPTION: SE         Gln Phe Trp Ala Thr Val Ser Tyr Tyr Glu Leu         5       10         Gly Glu Gln Val Lys Val Ser Ser Thr Thr Ile       20       25         Phe Thr Asp Pro Cys Ile Asn Gly Ser Lys Ile       35       40         Phe Ser Asn Val Asn Arg Asn Ala Thr Ile Glu 50       55         His Ile Gly Asn Gly Val Lys Leu Thr Tyr Val 65       55         Ser Leu Phe Ala Gln Cys Glu Ser Asp Ser Ala 80         Ser Leu Phe Ala Glu Cys Glu Ser Asp Gly Phe His 95         Nam Tyr Ile Asn Gly Phe His 100         100         Val Lys Ile Arg Gln Leu Glu Asp Cys Ser Leu Lys Ile 110         Ile Phe Arg Gln Leu Glu Lys Glu Asp Cys Ser Arg 130         Ala Ser Phe Asp Gly Typ Glu Arg Glu Lys Cys Glu Lys Glu Arg Glu L</td><td>(v)         FRAGMENT TYPE: internal           (vii)         ORIGINAL SOURCE:           (viii)         POSITION IN GENOME:           (viii)         POSITION IN GENOME:           (ix)         FEATURE:           (xi)         SEQUENCE DESCRIPTION:         SEQ ID           Gln         Phe         Trp         Ala         Thr         Val         Leu         Asa         SEQ ID           Gln         Phe         Trp         Ala         Thr         Val         Leu         Asa         ESEQ ID           Gln         Phe         Asa         SEQ ID           Gln         Asa         SEQ ID           Asa         <th co<="" td=""><td>(vi) FRAGMENT TYPE: internal           (vii) ORIGINAL SOURCE:           (viii) POSITION IN GENOME:           (ix) FEATURE:           (xi) SEQUENCE DESCRIPTION:         SEQ ID NO.:           GIN Pine Trp Ala Thr Val Ser Tyr Tyr Glu Leu Asn Thr 5 10           GIN GIN Val Lys Val Ser Ser Thr Thr Thr Thr Thr Thr Thr Thr Thr Th</td><td>(v) FRAGMENT TYPE: internal (vi) ORIGINAL SOURCE: (viii) IMMEDIATE SOURCE: (viii) POSITION IN GENOME: (ix) FEATURE: (x) PUBLICATION INFORMATION: (xi) SEQUENCE DESCRIPTION: (xi) SEQUEN</td></th></td></th>	<td>  (vi</td> <td>  (v)   FRAGMENT TYPE: into</td> <td> </td> <td>(v) FRAGMENT TYPE: internal         (vi) ORIGINAL SOURCE:         (vii) IMMEDIATE SOURCE:         (viii) POSITION IN GENOME:         (ix) FEATURE:         (xi) SEQUENCE DESCRIPTION: SE         Gln Phe Trp Ala Thr Val Ser Tyr Tyr Glu Leu         5       10         Gly Glu Gln Val Lys Val Ser Ser Thr Thr Ile       20       25         Phe Thr Asp Pro Cys Ile Asn Gly Ser Lys Ile       35       40         Phe Ser Asn Val Asn Arg Asn Ala Thr Ile Glu 50       55         His Ile Gly Asn Gly Val Lys Leu Thr Tyr Val 65       55         Ser Leu Phe Ala Gln Cys Glu Ser Asp Ser Ala 80         Ser Leu Phe Ala Glu Cys Glu Ser Asp Gly Phe His 95         Nam Tyr Ile Asn Gly Phe His 100         100         Val Lys Ile Arg Gln Leu Glu Asp Cys Ser Leu Lys Ile 110         Ile Phe Arg Gln Leu Glu Lys Glu Asp Cys Ser Arg 130         Ala Ser Phe Asp Gly Typ Glu Arg Glu Lys Cys Glu Lys Glu Arg Glu L</td> <td>(v)         FRAGMENT TYPE: internal           (vii)         ORIGINAL SOURCE:           (viii)         POSITION IN GENOME:           (viii)         POSITION IN GENOME:           (ix)         FEATURE:           (xi)         SEQUENCE DESCRIPTION:         SEQ ID           Gln         Phe         Trp         Ala         Thr         Val         Leu         Asa         SEQ ID           Gln         Phe         Trp         Ala         Thr         Val         Leu         Asa         ESEQ ID           Gln         Phe         Asa         SEQ ID           Gln         Asa         SEQ ID           Asa         <th co<="" td=""><td>(vi) FRAGMENT TYPE: internal           (vii) ORIGINAL SOURCE:           (viii) POSITION IN GENOME:           (ix) FEATURE:           (xi) SEQUENCE DESCRIPTION:         SEQ ID NO.:           GIN Pine Trp Ala Thr Val Ser Tyr Tyr Glu Leu Asn Thr 5 10           GIN GIN Val Lys Val Ser Ser Thr Thr Thr Thr Thr Thr Thr Thr Thr Th</td><td>(v) FRAGMENT TYPE: internal (vi) ORIGINAL SOURCE: (viii) IMMEDIATE SOURCE: (viii) POSITION IN GENOME: (ix) FEATURE: (x) PUBLICATION INFORMATION: (xi) SEQUENCE DESCRIPTION: (xi) SEQUEN</td></th></td>	(vi	(v)   FRAGMENT TYPE: into		(v) FRAGMENT TYPE: internal         (vi) ORIGINAL SOURCE:         (vii) IMMEDIATE SOURCE:         (viii) POSITION IN GENOME:         (ix) FEATURE:         (xi) SEQUENCE DESCRIPTION: SE         Gln Phe Trp Ala Thr Val Ser Tyr Tyr Glu Leu         5       10         Gly Glu Gln Val Lys Val Ser Ser Thr Thr Ile       20       25         Phe Thr Asp Pro Cys Ile Asn Gly Ser Lys Ile       35       40         Phe Ser Asn Val Asn Arg Asn Ala Thr Ile Glu 50       55         His Ile Gly Asn Gly Val Lys Leu Thr Tyr Val 65       55         Ser Leu Phe Ala Gln Cys Glu Ser Asp Ser Ala 80         Ser Leu Phe Ala Glu Cys Glu Ser Asp Gly Phe His 95         Nam Tyr Ile Asn Gly Phe His 100         100         Val Lys Ile Arg Gln Leu Glu Asp Cys Ser Leu Lys Ile 110         Ile Phe Arg Gln Leu Glu Lys Glu Asp Cys Ser Arg 130         Ala Ser Phe Asp Gly Typ Glu Arg Glu Lys Cys Glu Lys Glu Arg Glu L	(v)         FRAGMENT TYPE: internal           (vii)         ORIGINAL SOURCE:           (viii)         POSITION IN GENOME:           (viii)         POSITION IN GENOME:           (ix)         FEATURE:           (xi)         SEQUENCE DESCRIPTION:         SEQ ID           Gln         Phe         Trp         Ala         Thr         Val         Leu         Asa         SEQ ID           Gln         Phe         Trp         Ala         Thr         Val         Leu         Asa         ESEQ ID           Gln         Phe         Asa         SEQ ID           Gln         Asa         SEQ ID           Asa         Asa <th co<="" td=""><td>(vi) FRAGMENT TYPE: internal           (vii) ORIGINAL SOURCE:           (viii) POSITION IN GENOME:           (ix) FEATURE:           (xi) SEQUENCE DESCRIPTION:         SEQ ID NO.:           GIN Pine Trp Ala Thr Val Ser Tyr Tyr Glu Leu Asn Thr 5 10           GIN GIN Val Lys Val Ser Ser Thr Thr Thr Thr Thr Thr Thr Thr Thr Th</td><td>(v) FRAGMENT TYPE: internal (vi) ORIGINAL SOURCE: (viii) IMMEDIATE SOURCE: (viii) POSITION IN GENOME: (ix) FEATURE: (x) PUBLICATION INFORMATION: (xi) SEQUENCE DESCRIPTION: (xi) SEQUEN</td></th>	<td>(vi) FRAGMENT TYPE: internal           (vii) ORIGINAL SOURCE:           (viii) POSITION IN GENOME:           (ix) FEATURE:           (xi) SEQUENCE DESCRIPTION:         SEQ ID NO.:           GIN Pine Trp Ala Thr Val Ser Tyr Tyr Glu Leu Asn Thr 5 10           GIN GIN Val Lys Val Ser Ser Thr Thr Thr Thr Thr Thr Thr Thr Thr Th</td> <td>(v) FRAGMENT TYPE: internal (vi) ORIGINAL SOURCE: (viii) IMMEDIATE SOURCE: (viii) POSITION IN GENOME: (ix) FEATURE: (x) PUBLICATION INFORMATION: (xi) SEQUENCE DESCRIPTION: (xi) SEQUEN</td>	(vi) FRAGMENT TYPE: internal           (vii) ORIGINAL SOURCE:           (viii) POSITION IN GENOME:           (ix) FEATURE:           (xi) SEQUENCE DESCRIPTION:         SEQ ID NO.:           GIN Pine Trp Ala Thr Val Ser Tyr Tyr Glu Leu Asn Thr 5 10           GIN GIN Val Lys Val Ser Ser Thr Thr Thr Thr Thr Thr Thr Thr Thr Th	(v) FRAGMENT TYPE: internal (vi) ORIGINAL SOURCE: (viii) IMMEDIATE SOURCE: (viii) POSITION IN GENOME: (ix) FEATURE: (x) PUBLICATION INFORMATION: (xi) SEQUENCE DESCRIPTION: (xi) SEQUEN

Leu Asp Arg Val Leu Ser Thr Met Gly Pro Thr Pro Arg Pro Ile

185 190 195 Ser Ser Ile Ser (9) INFORMATION FOR SEQ ID NO: 8 SEQUENCE CHARACTERISTICS: (i) (A) LENGTH: 198 amino acids (B) TYPE: amino acid (C) STRANDEDNESS: 10 (D) TOPOLOGY: linear (ii) MOLECULE TYPE: (A) DESCRIPTION: peptide (iii) HYPOTHETICAL: no (iv) ANTISENSE: no 15 (v) FRAGMENT TYPE: internal (vi) ORIGINAL SOURCE: (vii) IMMEDIATE SOURCE: (viii) POSITION IN GENOME: (ix) FEATURE: 20 PUBLICATION INFORMATION: (x) (xi) SEQUENCE DESCRIPTION: SEQ ID NO.: 8 Lys Ser Trp Ala Gln Ile Thr Tyr Phe Glu Leu Asn Ser Arg Val 10 Gly Glu Val Phe Lys Leu Val Asn Leu Ser Ile Thr Val Asp Gly 25 20 25 30 Tyr Thr Asn Pro Ser Asn Ser Asn Thr Arg Ile Cys Leu Gly Gln 35 40 Leu Thr Asn Val Asn Arg Asn Gly Thr Ile Glu Asn Thr Arg Met 50 55 60 His Ile Gly Lys Gly Ile Gln Leu Asp Asn Lys Glu Asp Gln Met 30

70

75

	His	Ile	Met	Ile	Thr	Asn	Asn	Ser	Asp	Met	Pro	Val	Phe	Val	Glr
					80					85					90
	Ser	Lys	Asn	Thr	Asn	Leu	Met	Met	Asn	Met	Pro	Leu	Val	Lys	Val
					95					100					105
5	Cys	Arg	Ile	Pro	Pro	His	Ser	Gln	Leu	Cys	Val	Phe	Glu	Phe	Asn
					110					115					120
	Leu	Phe	Phe	Gln	Met	Leu	Glu	Gln	Ser	Cys	Asn	Asp	Ser	Asp	Gly
					125					130					135
	Leu	Asn	Glu	Leu	Ser	Lys	His	Cys	Phe	Ile	Arg	Ile	Ser	Phe	Val
10					140					145					150
	Lys	Gly	Trp	Gly	Glu	Asp	Tyr	Pro	Arg	Gln	Asp	Val	Thr	Ser	Thr
					155					160					165
	Pro	Cys	Trp	Leu	Glu	Leu	Arg	Leu	Asn	Val	Pro	Leu	Ala	Tyr	Ile
					170					175					180
15	Asp	Gln	Lys	Met		Gln	Thr	Pro	Arg	Thr	Asn	Leu	Met	Pro	Asn
					185					190					195
	Ser	Met	Thr												

#### 20 (10)INFORMATION FOR SEQ ID NO: 9

- (i) SEQUENCE CHARACTERISTICS
  - (A) LENGTH: 224 amino acids
  - (B) TYPE: amino acid
  - (C) STRANDEDNESS:
- 25 (D) TOPOLOGY: linear
  - (ii) MOLECULE TYPE:
    - (A) DESCRIPTION: peptide
  - (iii) HYPOTHETICAL: no
  - (iv) ANTISENSE: no
- 30 (v) FRAGMENT TYPE: internal
  - (vi) ORIGINAL SOURCE:
  - (vii) IMMEDIATE SOURCE:

## (viii) POSITION IN GENOME:

- (ix) FEATURE:
- (x) PUBLICATION INFORMATION:

			()	(i)	SEQU	JENC	E DE	SCRI	PTIO	N:	SE	QID	NO.:	9	
5	Leu	Asp	Asn	Trp	Cys	Ser	Ile	Ile	Tyr	Tyr	Glu	Leu	Asp	Thr	Pro
					5					10					15
	Ile	Gly	Glu	Thr	Phe	Lys	Val	Ser	Ala	Arg	Asp	His	Gly	Lys	Val
					20					25					30
	Ile	Val	Asp	Gly	Gly	Met	Asp	Pro	His	Gly	Glu	Asn	Glu	Gly	Arg
10					35					40					45
	Leu	Cys	Leu	Gly	Ala	Leu	Ser	Asn	Val	His	Arg	Thr	Glu	Ala	Ser
					50					55					60
	Glu	Lys	Ala	Arg	Ile	His	Ile	Gly	Arg	Gly	Val	Glu	Leu	Thr	Ala
					65					70					75
15	His	Ala	Asp	Gly	Asn	Ile	Ser	Ile	Thr		Asn	Cys	Lys	Ile	Phe
					80					85					90
	Val	Arg	Ser	Gly	Tyr	Leu	Asp	Tyr	Thr		Gly	Ser	Glu	Tyr	
	_	_			95	_,		_	_	100	_				105
20	Ser	Lys	Ala	His	Arg	Phe	Thr	Pro	Asn		Ser	Ser	Phe	Thr	
20	701	<b>-</b>	<b>T</b> 3 -	3	110	37 -	<b>.</b>	<b>3</b> 7 4-	<b>~</b> 1	115	_		_	~	120
	Pne	Asp	TIE	Arg	Trp 125	Ala	ıyr	met	GIn		Leu	Arg	Arg	Ser	-
	) an	Cox	7.00	C1		170 J	7	71-	G1-	130	7.1.	7.1 -	77- 7	71-	135
	Asp	SET	ASII	Giu	Ala 140	vai	Arg	Ala	GIII	145	Ата	Ala	vai	Ата	150
25	ጥነተ	Δla	Pro	Mat	Ser	172 l	Mot	Dro	בות		Mot	Dro	50×	Cor	
23		2324	110	1100	155	Val	1100	110	niu	160	Mec	FIO	361	361	165
	Val	Asp	Ara	Met	Arg	Ara	Asp	Phe	Cvs		Tle	Ala	Tle	Ser	
			· 3		170	5			-1-	175				501	180
	Val	Lys	Ala	Trp	Gly	Asp	Val	Tyr	Gln		Lvs	Thr	Ile	Lvs	
30		_		-	185	_		-		190	•				195
	Thr	Pro	Cys	Trp	Ile	Glu	Val	Thr	Leu	His	Arg	Pro	Leu	Gln	
					200					205					210
	Leu	Asp	Gln	Leu	Leu	Lys	Asn	Ser	Ser	Gln	Phe	Gly	Ser	Ser	
					215					220					

	(11)	)INF(	ORM.	ATIC	N FC	OR SE	EQ ID	NO:	10						
			(j	i)	SEQU	JENC	E CF	IARA	CTE	RISTI	CS:				
					(A)	LE	NGT	H: 5	0 am	ino a	cids				
5					(B)	TY	PE:	amin	o aci	d					
					(C)	STF	RANE	EDN	ESS:						
					(D)	TO	POLO	OGY:	line	ar					
			(i	i)	MOL	ECUI	ETY	PE:							
					(A)	DE	SCRI	PTIC	N: 1	peptic	ie				
10			(i	ii)	HYP	OTH	ETIC	AL: 1	no						
			(i	v)	ANT	ISEN	SE:	no							
			( '	v)	FRA	GME	NT I	YPE	: int	ernal					
			()	vi)	ORIC	INAI	LSO	JRCE	<b>:</b> :						
			()	vii)	IMM!	EDIA	TE S	OUR	CE:						
15			7)	viii)	POSI	TION	IN	GENC	ME:						
			(i	x)	FEAT	URE	:								
			()	x)	PUBI	LICA	TION	INF	ORM	ATIC	N:				
			()	(i)	SEQU	JENC	E DE	SCRI	PTIO	N:	SE	Q ID	NO.:	10	
<b>3</b> 0	Phe	Val	Lys	Gly	Trp	Gly	Ala	Glu	Tyr		Arg	Gln	Thr	Val	
20	Ser	Thr	Pro	Cvs	5 Trp	Tle	Glu	I.e.u	Hic	10 Leu	Hic	Gly	Dro	Lou	15
	DCI	1111	110	Cys	20	110	GIU	Deu	1113	25	1113	GIY	FIO	Deu	30
	Trp	Leu	Asp	Lys	Val	Leu	Thr	Gln	Met	Gly	Ser	Pro	Ser	Val	Arg
					35					40					45
25	Суѕ	Ser	Ser	Met	Ser 50										
					30										
	(12)	INF	ORM.	ATIC	N FO	R SE	Q ID	NO:	11						
			(i	.)	SEQU	JENC	E CH	[ARA	CTER	ISTIC	CS:				
30					(A)	LE	\GTI	H: 50	0 am	ino a	cids				
					(B)	TY	PE:	amin	o aci	d					

					(C)	ST	RANI	DEDN	IESS:						
					(D)	TO	POL	OGY	: line	ar					
			(	ii)	MOL	ECU	LET	YPE:							
					(A)	DE	SCR	IPTIC	N:	pepti	de				
5			(i	iii)	HYP	отн	ETIC	AL:	no						
			<b>(</b> i	iv)	ANT	ISEN	ISE:	no							
			(	v)	FRA	GME	NT T	ГҮРЕ	: int	ernal					
			(	vi)	ORIC	3INA	L SO	URCE	Ē:						
			()	vii)	IMM	EDLA	TE S	OUR	CE:						
10			()	viii)	POSI	TION	IN (	GENC	OME:						
			(i	x)	FEA7	TURE	:								
			()	<b>(</b> )	PUB	LICA	TION	INF	ORM	IATIO	ON:				
			()	(i)	SEQU	JENC	EDE	SCRI	PTIO	N:	SE	Q ID	NO.:	11	
	Phe	Val	Lys	Gly	Trp	Gly	Ala	Glu	Tyr	Arg					Thr
15	_		_	_	5					10					15
	Ser	Thr	Pro	Cys	Trp 20	Ile	Glu	Leu	His		His	Gly	Pro	Leu	
	Trp	Leu	Asp	Lvs	-	Leu	Thr	Gln	Met	25 Gly	Sor	Dro	Sor	Tlo	30
			_	-	35					40	501	110	261	TIE	45
20	Cys	Ser	Ser	Met	Ser										
					50										
	(13)	INFO	ORM <i>A</i>	OITA	N FO	R SE	QЮ	NO:	12						
			(i)	)	SEQL	JENC.	E CH	ARA	TER	ISTIC	CS:				
25					(A)			I: 50							
					(B)	TYF	E: a	amino	acio	i					
				(	(C)	STR	AND	EDNE	ESS:						
				(	(D)	TOP	OLO	GY:	linea	r					
			(ii	) ]	MOLE	ECUL	ETY.	PE:							
30				(	(A)	DES	CRIE	OIT	V: p	eptid	е				
			(ii	i) l	HYPC				_	•					

			(i	v)	ANT	ISEN	SE:	no							
			()	v)	FRA	GME	NT T	YPE	inte	ernal					
			()	vi)	ORIC	INAI	SOU	JRCE	<b>:</b> :						
			()	vii)	IMM	EDIA	TE S	OUR	CE:						
5			()	viii)	POSI	TION	INC	GENC	ME:						
			(i	x)	FEA1	TURE	:								
			()	<b>(</b> )	PUBI	LICA	TION	INF	ORM	ATIO	ON:				
			()	(i)	SEQU	JENC	EDE	SCRI	PTIO	N:	SE	Q ID	NO.:	12	
	Phe	Val	Lys	Gly	Trp	Gly	Ala	Glu	Tyr	His	Arg	Gln	Asp	Val	Thr
10					5					10					15
	Ser	Thr	Pro	Cys	Trp 20	Ile	Glu	Leu	His	Leu 25	His	Gly	Pro	Leu	
	Trp	Leu	Asp	Lvs	Val	Leu	Thr	Gln	Met		Ser	Pro	His	Asn	30 Ala
			<b>-</b>		35					40				11011	45
15	Ile	Ser	Ser	Met	Ser										
					50										
	(1.4)	INIE	<b>7DM</b>	A TIC	NI EC	n er	O ID	NO.	12						
20	(14)	)INF			N FO		_			וכייו	TC.				
20			(i	.)	SEQU										
					(A)			1: 50			cias				
					(B)			amino		1					
					(C)			EDNI		_					
25			<i>(</i> :	: \	(D)			OGY:	mea	ır					
25			(1	i)	MOL				NI. ~		la.				
			<i>(</i> :	::1	(A) HYP				_	ерис	ie				
			-	,	ANT				10						
			Ţ												
20			,	•	FRAC					rnal					
30				•	ORIC										
			(1	/ii)	IMM!	CUIA	TE 20	OUK	LE:						

(viii) POSITION IN GENOME:

(ix) FEATURE: (x) PUBLICATION INFORMATION: (xi) SEQUENCE DESCRIPTION: SEQ ID NO.: 13 Phe Val Lys Gly Trp Gly Ala Glu Tyr His Arg Gln Asp Val Thr 5 10 15 Ser Thr Pro Cys Trp Ile Glu Leu His Leu His Gly Pro Leu Gln 20 25 Trp Leu Asp Lys Val Leu Thr Gln Met Gly Ser Pro His Asn Pro 10 35 40 45 Ile Ser Ser Met Ser 50 15 (15)INFORMATION FOR SEQ ID NO: 14 (i) SEQUENCE CHARACTERISTICS: (A) LENGTH: 50 amino acids (B) TYPE: amino acid (C) STRANDEDNESS: 20 (D) TOPOLOGY: linear (ii) MOLECULE TYPE: (A) DESCRIPTION: peptide (iii) HYPOTHETICAL: no (iv) ANTISENSE: no 25 FRAGMENT TYPE: internal (v) (vi) ORIGINAL SOURCE: (vii) IMMEDIATE SOURCE: (viii) POSITION IN GENOME: (ix) FEATURE: 30 PUBLICATION INFORMATION: (x)

			()	κi)	SEQU	JENC	E DE	SCRI	PTIO	N:	SE	Q ID	NO.:	14	
	Phe	Val	Lys	Gly	Trp	Gly	Ala	Glu	Tyr	His	Arg	Gln	Asp	Val	Thr
					5					10					15
	Ser	Thr	Pro	Cys	Trp	Ile	Glu	Leu	His	Leu	His	Gly	Pro	Leu	Glr
5					20					25					30
	Trp	Leu	Asp	Lys	Val	Leu	Thr	Gln	Met		Ser	Pro	Leu	Asn	
	<b>-</b> 1 -	G	0	<b>\</b>	35					40					45
	TTE	Ser	Ser	Met	50										
10					50										
	(16)	INFO	ORM.	ATIC	N FC	R SE	O ID	NO:	15						
	(- /		(i		SEQU		-			USTIC	CS:				
			`	,	(A)		<b>VGT</b> I								
15					(B)		PE:								
					(C)										
					(D)					ar.					
			(i	i)	MOL										
			\-	· <b>-</b> /	(A)		SCRI		N∙ r	nentid	le				
20			(i	ii)	HYP				•	орис					
20			•	•	ANT				10						
			•	•	FRA				. :_+.						
			`	•						ernai					
			•	-	ORIC										
			•	·	IMM										
25			(7	viii)	POSI	TION	INC	BENC	ME:						
			(i	x)	FEA]	TURE	:								
			()	x)	PUB	LICA	TION	INF	ORM	ATIC	N:				
			()	xi)	SEQU	JENC	E DE	SCRI	PTIO	N:	SE	Q ID	NO.:	15	
	Phe	Val	Lys	Gly	Trp	Gly	Pro	Asp	Tyr	Pro	Arg	Gln	Ser	Ile	Lys
30	_				5					10			_		15
	Glu	Thr	Pro	Cys	Trp	Ile	Glu	Leu	His	Leu 25	His	Arg A	Ala I	Jeu G	ln 30
					2.11					17					< (1

Leu Leu Asp Glu Val Leu His Thr Met Pro Ile Ala Asp Pro Gln

35 40 45 Pro Leu Asp (17) INFORMATION FOR SEQ ID NO: 16 5 SEQUENCE CHARACTERISTICS: (i) (A) LENGTH: 29 amino acids (B) TYPE: amino acid (C) STRANDEDNESS: 10 (D) TOPOLOGY: linear MOLECULE TYPE: (ii) (A) DESCRIPTION: peptide (iii) HYPOTHETICAL: no (iv) ANTISENSE: no (v) FRAGMENT TYPE: internal 15 (vi) ORIGINAL SOURCE: (vii) IMMEDIATE SOURCE: (viii) POSITION IN GENOME: (ix) FEATURE: 20 (x) PUBLICATION INFORMATION: (xi) SEQUENCE DESCRIPTION: SEQ ID NO.: 16 Phe Ala Lys Gly Trp Gly Pro Cys Tyr Ser Arg Gln Phe Ile Thr 5 10 15 Ser Cys Pro Cys Trp Leu Glu Ile Leu Leu Asn Asn Pro Arg 25 20 25 (18) INFORMATION FOR SEQ ID NO: 17 (i) **SEQUENCE CHARACTERISTICS:** (A) LENGTH: 28 amino acids 30 (B) TYPE: amino acid

		(C) STRANDEDNESS:
		(D) TOPOLOGY: linear
	(ii)	MOLECULE TYPE:
		(A) DESCRIPTION: peptide
5	(iii)	HYPOTHETICAL: no
	(iv)	ANTISENSE: no
	(v)	FRAGMENT TYPE: internal
	(vi)	ORIGINAL SOURCE:
	(vii)	IMMEDIATE SOURCE:
10	(viii)	POSITION IN GENOME:
	(ix)	FEATURE:
	(x)	PUBLICATION INFORMATION:
	(xi)	SEQUENCE DESCRIPTION: SEQ ID NO.: 17
1.5	Phe Val Lys Gly	Trp Gly Gln Cys Tyr Thr Arg Gln Phe Ile Ser
15	Ser Cvs Pro Cvs	5 10 15 Trp Leu Glu Val Ile Phe Asn Ser Arg
		20 25
	(19)INFORMATION	ON FOR SEQ ID NO: 18
20	(i)	SEQUENCE CHARACTERISTICS:
		(A) LENGTH: 25 amino acids
		(B) TYPE: amino acid
		(C) STRANDEDNESS:
		(D) TOPOLOGY: linear
25	(ii)	MOLECULE TYPE:
		(A) DESCRIPTION: peptide
	(iii)	HYPOTHETICAL: no
	(iv)	ANTISENSE: no
	(v)	FRAGMENT TYPE: internal
30	(vi)	ORIGINAL SOURCE:

	(vii)	IMMEDIATE SOURCE:							
	(viii	POSITION IN GENOME:							
	(ix)	FEATURE:							
	(x)	PUBLICATION INFORMATION:							
5	(xi)	SEQUENCE DESCRIPTION: SEQ ID NO.: 18							
	Leu Gly Phe Il	e Ala Ala Asp Asn Lys Asp Asn Gly Thr Trp Th							
	Gln Leu Trp Le	u Val Ser Asp Tyr His Glu							
		20 25							
10									
	(20)INFORMATI	ON FOR SEQ ID NO: 19							
	(i)	SEQUENCE CHARACTERISTICS:							
		(A) LENGTH: 25 amino acids							
	(B) TYPE: amino acid								
15		(C) STRANDEDNESS:							
		(D) TOPOLOGY: linear							
	(ii)	MOLECULE TYPE:							
		(A) DESCRIPTION: peptide							
	(iii)	HYPOTHETICAL: no							
20	(iv)	ANTISENSE: no							
	(v)	FRAGMENT TYPE: internal							
	(vi)	ORIGINAL SOURCE:							
	(vii)	IMMEDIATE SOURCE:							
	(viii)	POSITION IN GENOME:							
25	(ix)	FEATURE:							
	(x)	PUBLICATION INFORMATION:							
	(xi)	SEQUENCE DESCRIPTION: SEQ ID NO.: 19							
	Leu Gly Phe Ile	e Ala Ala Asp Ile Lys Gly Thr Gly Thr Trp Thr							
30		5 10 15							

Gln Leu Tyr Leu Ile Thr Asp Tyr His Glu
20 25

#### (21) INFORMATION FOR SEQ ID NO: 20

5 (i) SEQUENCE CHARACTERISTICS:

- (A) LENGTH: 25 amino acids
- (B) TYPE: amino acid
- (C) STRANDEDNESS:
- (D) TOPOLOGY: linear
- 10 (ii) MOLECULE TYPE:
  - (A) DESCRIPTION: peptide
  - (iii) HYPOTHETICAL: no
  - (iv) ANTISENSE: no
  - (v) FRAGMENT TYPE: internal
- 15 (vi) ORIGINAL SOURCE:
  - (vii) IMMEDIATE SOURCE:
  - (viii) POSITION IN GENOME:
  - (ix) FEATURE:
  - (x) PUBLICATION INFORMATION:
- 20 (xi) SEQUENCE DESCRIPTION: SEQ ID NO.: 20

Leu Gly Phe Ile Ala Ala Asp Ile Lys Gly Asn Gly Thr Trp Thr
5 10 15

Gln Met Leu Leu Ile Thr Asp Tyr His Glu 20 25

25

#### (22) INFORMATION FOR SEQ ID NO: 21

- (i) SEQUENCE CHARACTERISTICS:
  - (A) LENGTH: 25 amino acids
  - (B) TYPE: amino acid
- 30 (C) STRANDEDNESS:

		(D) TOPOLOGY: linear
	(ii)	MOLECULE TYPE:
		(A) DESCRIPTION: peptide
	(iii)	HYPOTHETICAL: no
5	(iv)	ANTISENSE: no
	(v)	FRAGMENT TYPE: internal
	(vi)	ORIGINAL SOURCE:
	(vii)	IMMEDIATE SOURCE:
	(viii)	POSITION IN GENOME:
10	(ix)	FEATURE:
	(x)	PUBLICATION INFORMATION:
	(xi)	SEQUENCE DESCRIPTION: SEQ ID NO.: 21
	Leu Gly Phe Ile	e Ala Ser Asp Met Thr Ser Arg Asn Ser Ser Thr
15	Olm Lou Mam Lou	5 10 15
13	GIN Led TIP Led	I Ile Thr His Tyr His Glu 20 25
	(23)INFORMATION	ON FOR SEQ ID NO: 22
	(i)	SEQUENCE CHARACTERISTICS:
20		(A) LENGTH: 25 amino acids
		(B) TYPE: amino acid
		(C) STRANDEDNESS:
		(D) TOPOLOGY: linear
	(ii)	MOLECULE TYPE:
25		(A) DESCRIPTION: peptide
	(iii)	HYPOTHETICAL: no
	(iv)	ANTISENSE: no
	(v)	FRAGMENT TYPE: internal
	(vi)	ORIGINAL SOURCE:
30	(vii)	IMMEDIATE SOURCE:

	(viii	) POSITION IN GENOME:
	(ix)	FEATURE:
	(x)	PUBLICATION INFORMATION:
	(xi)	SEQUENCE DESCRIPTION: SEQ ID NO.: 22
5	Leu Gly Phe Il	e Ala Ser Asp Met Thr Ser Arg His Ser Ser Thr 5 10 15
	Gln Leu Trp Le	u Ile Thr His Tyr His Glu 20 25
10	(24)INFORMATI	ION FOR SEQ ID NO: 23
	(i)	SEQUENCE CHARACTERISTICS:
		(A) LENGTH: 25 amino acids
		(B) TYPE: amino acid
		(C) STRANDEDNESS:
15		(D) TOPOLOGY: linear
	(ii)	MOLECULE TYPE:
		(A) DESCRIPTION: peptide
	(iii)	HYPOTHETICAL: no
	(iv)	ANTISENSE: no
20	(v)	FRAGMENT TYPE: internal
	(vi)	ORIGINAL SOURCE:
	(vii)	IMMEDIATE SOURCE:
	(viii)	POSITION IN GENOME:
	(ix)	FEATURE:
25	(x)	PUBLICATION INFORMATION:
	(xi)	SEQUENCE DESCRIPTION: SEQ ID NO.: 23
	Leu Gly Phe Ile	e Gly Ser Asp Met Thr Ser Arg Asn Ser Cys Thr
	Gln Leu Tro Lo	5 10 15 1 Met Thr His Tyr Tyr Pro
30	orn ned ith ned	20 25
		- <del></del>

	(25)INFORMATION FOR SEQ ID NO: 24														
			<b>(</b> i	i)	SEQ	JENC	CE CH	IARA	CTE	RISTI	CS:				
					(A)	LE	NGT	H: 1	99 aı	mino	acid	S			
5					(B)	TY	PE:	amin	o aci	d					
					(C)	STE	RAND	EDN	ESS:						
					(D)	TO	POLO	OGY:	line	ar			-		
			(i	i)	MOL	ECUI	LETY	PE:							
					(A)	DE	SCRI	PTIC	N: 1	peptio	ie				
10			(i	ii)	HYP	отні	ETIC.	AL: 1	no						
			(i	v)	ANT	ISEN	SE:	no							
			(,	v)	FRA	GME	NT I	YPE	: inte	ernal					
			()	vi)	ORIC	INAI	LSOU	JRCE	<b>:</b> :						
			('	vii)	IMM	EDIA	TE S	OUR	CE:						
15			()	viii)	POSI	TION	IN C	SENC	ME:						
			(i	x)	FEA7	TURE	:								
			()	<b>(</b> )	PUBI	LICA	TION	INF	ORM	ATIO	ON:				
			()	(i)	SEQU	JENC	E DE	SCRI	PTIO	N:	SE	Q ID	NO.:	24	
	Glu	Pro	Lys	His	Trp	Cys	Ser	Ile	Val	Tyr	Tyr	Glu	Leu	Asn	Asn
20					5					10					15
	Arg	Val	Gly	Glu	Ala	Phe	His	Ala	Ser		Thr	Ser	Val	Leu	
	Asp	Glv	Phe	Thr	20 Asp	Pro	Ser	Asn	Asn	25 Lvs	Asn	Ara	Phe	Cve	30 Len
	<i>F</i>	4			35		-			40	11011	**** 9	1110	Cys	45
25	Gly	Leu	Leu	Ser	Asn	Val	Asn	Arg	Asn	Ser	Thr	Ile	Glu	Asn	Thr
					50					55					60
	Arg	Arg	His	Ile	Gly	Lys	Gly	Val	His		Tyr	Tyr	Val	Gly	
	Glu	Val	Tvr	Ala	65 Glu	Cvs	ī,en	Ser	Asp	70 Ser	Ser	Tle	Phe	V=1	75 Gln
30			-1-		80	<b>-</b> 75	204	201	nsp		DCI	115	1116	vaı	GIII

	Ser	Arg	Asn	Cys	Asn 95	Tyr	His	His	Gly	Phe	His	Pro	Thr	Thr	Val
	Cvs	Lvs	Ile	Pro		Glv	Cvs	Ser	Len		Tle	Phe	Asn	Asn	
	CyD	ביינם			110	0_1	O <sub>I</sub> O	501	Deu	115			11011		120
5	Glu	Phe	Ala	Gln	Leu	Leu	Ala	Gln	Ser	Val	Asn	His	Gly	Phe	Glu
					125					130					135
	Thr	Val	Tyr	Glu	Leu	Thr	Lys	Met	Cys	Thr	Ile	Arg	Met	Ser	Phe
					140					145					150
	Val	Lys	Gly	Trp		Ala	Glu	Tyr	His		Gln	Asp	Val	Thr	
10	mb	D	O a	Шэгэ	155	C1	<b>T</b> 1.	774 -	T 011	160	~1·-	Desa	T	Q1 m	165
	Thr	Pro	Cys	Trp	170	GIU	TIE	nis	ьeu	175	GIA	PIO	ьeu	GIII	180
	Leu	Asp	Lys	Val		Thr	Gln	Met	Gly		Pro	His	Asn	Pro	
		-			185				_	190					195
15	Ser	Ser	Val	Ser											
	(26)	)INF(	ORM	ATIC	N FC	R SE	Q ID	NO:	25						
			(i	)	SEQU	JENC	E CH	ARA	CTER	USTIC	CS:				
					(A)	LEI	NGTI	H: 1	98 ar	nino	acids				
20					(B)	TY	PE:	amin	o aci	d					
					(C)	STF	RAND	EDN	ESS:						
					(D)	TO	POLO	OGY:	linea	ar					
			(i	i)	MOL	ECUI	ETY	PE:							
					(A)	DE	SCRI	PTIO	N: ŗ	peptic	le				
25			(i	iii)	HYP	OTH	ETIC.	AL: 1	10	-					
			(i	iv)	ANT	ISEN	SE:	no							
			-	v)	FRA	GME	NT T	YPE	: inte	ernal					
			•	vi)	ORIC	JINAI	LSOU	JRCE							
			•	•	IMM	EDIA	TE S	OUR	CE:						
30			•	•	POSI										
-			•	ŕ	FEAT										

## (x) PUBLICATION INFORMATION:

			(xi) SEQUENCE DESCRIPTION:						SEQ ID NO.: 25						
	Glu	Pro	Ala	Phe	Trp	Cys	Ser	Ile	Ala	Tyr	Tyr	Glu	Leu	Asn	Gln
					5					10					15
5	Arg	Val	Gly	Glu	Thr	Phe	His	Ala	Ser	Gln	Pro	Ser	Leu	Thr	Val
					20					25					30
	Asp	Gly	Phe	Thr	Asp	Pro	Ser	Asn	Ser	Glu	Arg	Phe	Cys	Leu	Gly
					35					40					45
	Leu	Leu	Ser	Asn	Val	Asn	Arg	Asn	Ala	Thr	Val	Glu	Met	Thr	Arg
10					50					55					60
	Arg	His	Ile	Gly	Arg	Gly	Val	Arg	Leu	Tyr	Tyr	Ile	Gly	Gly	Glu
					65					70					75
	Val	Phe	Ala	Glu	Суs	Leu	Ser	Asp	Ser	Ala	Ile	Phe	Val	Gln	Ser
					80					85					90
15	Pro	Asn	Cys	Asn	Gln	Arg	Tyr	Gly	Trp	His	Pro	Ala	Thr	Val	Cys
					95					100					105
	Lys	Ile	Pro	Pro	_	Cys	Asn	Leu	Lys		Phe	Asn	Asn	Gln	
					110					115					120
	Phe	Ala	Ala	Leu		Ala	Gln	Ser	Val		Gln	Gly	Phe	Glu	
20					125					130					135
	Val	Tyr	Gln	Leu		Arg	Met	Cys	Thr		Arg	Met	Ser	Phe	
					140					145		_			150
	Lys	Gly	Trp	Gly		Glu	Tyr	Arg	Arg		Thr	Val	Thr	Ser	
~ ~		_	_		155	_		_		160		_	-1	_	165
25	Pro	Суѕ	Trp	Ile		Leu	Hıs	Leu	His		Pro	Leu	Gln	Trp	
		_	7	_	170	~1		<b>~</b> 3	_	175	_	1	_	~	180
	Asp	Lys	Val	Leu		Gin	Met	GIY	Ser		Ser	Val	Arg	Cys	
	_				185					190					195
	Ser	Met	Ser												

#### WHAT IS CLAIMED IS:

1. A method of testing compounds, comprising the steps of:

- 5 a) providing (i) a Smad4 polypeptide comprising the L3 loop region, (ii) a complementary Smad polypeptide, and (iii) a compound to be tested; and
  - (b) contacting said Smad4 polypeptide with said complementary Smad polypeptide under conditions where binding can take place, wherein said contacting is performed in the presence and absence of said compound; and
  - c) detecting an increase or decrease in binding of said Smad4 polypeptide to said complementary Smad polypeptide in the presence of said compound.

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- 2. The method of claim 1, wherein the complementary Smad polypeptide is selected from the group consisting of Smad1, Smad2, Smad3, Smad5 and Smad8.
- 3. The method of claim 1, wherein said compound may be used to treat ailments selected from the group consisting of pancreatic cancer, breast cancer, ovarian cancer, colon cancer, esophageal cancer, head and neck cancers, fibrosis of the kidney, fibrosis of the liver, fibrosis of the lung, Alzheimer's disease, memory loss, inflammation, wound healing, bone growth, immunoregulation, blood cell formation and atherosclerosis.

4. The method of claim 1, wherein said detection is selected from the group consisting of solid support immobilization of one or the other Smad polypeptides, labeling of one or the other Smad polypeptides, scintillation proximity, homogeneous time resolved fluorescence, fluorescence resonance energy transfer and fluorescence polarization.

- 5. A method of testing compounds, comprising the 10 steps of:
  - a) providing (i) two Smad polypeptides from the same Smad family comprising the C-terminal domains of each, and (ii) a compound to be tested; and
- b) contacting said Smad polypeptides under conditions

  where binding can take place, wherein said contacting is performed
  in the presence and absence of said compound; and
  - c) detecting an increase or decrease in binding of said Smad polypeptides to each other in the presence of said compound.
- 5. The method of claim 5, wherein the families of Smad polypeptides are selected from the group consisting of Smad1, Smad2, Smad3, Smad4, Smad5, Smad6, Smad7 and Smad8.
- 7. The method of claim 5, wherein said drug may be used to treat ailments selected from the group consisting of pancreatic cancer, breast cancer, ovarian cancer, colon cancer.

esophageal cancer, head and neck cancers, fibrosis of the kidney, fibrosis of the liver, fibrosis of the lung, Alzheimer's disease, memory loss, inflammation, wound healing, bonc growth, immunoregulation, blood cell formation and atherosclerosis.

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- 8. The method of claim 5, wherein said detection is selected from the group consisting of solid support immobilization of one or the other Smad polypeptides, labeling of one or the other Smad polypeptides, scintillation proximity, homogeneous time resolved fluorescence, fluorescence resonance energy transfer and fluorescence polarization.
- 9. A method of testing compounds, comprising the steps of:
- a) providing (i) a Smad polypeptide comprising the C-terminal domain, (ii) a polypeptide comprising the L45 loop of the kinase domain corresponding to a receptor of the TGF-β or BMP family, and (iii) a test compound; and
- b) contacting said Smad polypeptide with said
  20 receptor polypeptide under conditions where phosphorylation can
  take place, wherein said contacting is performed in the presence and
  absence of said compound; and
- c) detecting an increase or decrease in the phosphorylation of said Smad polypeptide in the presence of said compound.

10. The method of claim 9, wherein the Smad polypeptide is selected from the group consisting of Smad1, Smad2, Smad3, Smad5 and Smad8.

- 11. The method of claim 9, wherein said drug may be used to treat ailments selected from the group consisting of pancreatic cancer, breast cancer, ovarian cancer, colon cancer, esophageal cancer, head and neck cancers, fibrosis of the kidney, fibrosis of the liver, fibrosis of the lung, Alzheimer's disease, memory loss, inflammation, wound healing, bone growth, immunoregulation, blood cell formation and atherosclerosis.
- 12. The method of claim 9, wherein said detection is selected from the group consisting of gel electrophoresis and scintillation counting.
  - 13. A method of testing compounds, comprising the steps of:
- a) providing (i) a Smad polypeptide comprising the α 20 helix 2 of the MH2 domain, (ii) a DNA binding polypeptide, and (iii) a compound to be tested; and
  - b) contacting said Smad polypeptide with said DNA binding polypeptide under conditions where binding can take place, wherein said contacting is performed in the presence and absence of said compound; and

c) detecting whether there is an increase in binding of said Smad polypeptide to said DNA binding polypeptide in the presence of said compound.

- 5 14. The method of claim 13, wherein the Smad polypeptide is selected from the group consisting of Smad1, Smad2, Smad3, Smad4, Smad5 and Smad8.
- 15. The method of claim 13, wherein said drug may be used to treat ailments selected from the group consisting of pancreatic cancer, breast cancer, ovarian cancer, colon cancer, esophageal cancer, head and neck cancers, fibrosis of the kidney, fibrosis of the liver, fibrosis of the lung, Alzheimer's disease, memory loss, inflammation, wound healing, bone growth, immunoregulation, blood cell formation and atherosclerosis.
  - 16. The method of claim 13, wherein said DNA binding polypeptide is selected from the group consisting of FAST1 and homologues of FAST1.

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17. The method of claim 13, wherein said detection is selected from the group consisting of solid support immobilization of one or the other Smad polypeptides, labeling of one or the other Smad polypeptides, scintillation proximity, homogeneous time resolved fluorescence, fluorescence resonance energy transfer and fluorescence polarization.

18. A method of testing compounds, comprising the steps of:

- a) providing (i) two Smad polypeptides comprising the C-terminus of each, (ii) a Smad polypeptide comprising the N-terminal domain, and (iii) a compound to be tested; and
  - b) contacting said Smad C-terminus polypeptides in the presence of said Smad N-terminal domain under conditions where binding can take place, wherein said contacting is performed in the presence and absence of said compound;
  - c) detecting whether there is an increase or decrease in binding of said Smad C-terminus domains in the presence of said compound due to inhibition of the autoinhibitory function of the N-terminal domain by said compound.

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- 19. The method of claim 18, wherein the Smad polypeptide is selected from the group consisting of Smad1, Smad2, Smad3, Smad4, Smad5 and Smad8.
- 20. The method of claim 18, wherein said drug may be used to treat ailments selected from the group consisting of pancreatic cancer, breast cancer, ovarian cancer, colon cancer, esophageal cancer, head and neck cancers, fibrosis of the kidney, fibrosis of the liver, fibrosis of the lung, Alzheimer's disease, memory loss, inflammation, wound healing, bone growth, immunoregulation, blood cell formation and atherosclerosis.

21. The method of claim 18, wherein said detection is selected from the group consisting of solid support immobilization of one or the other Smad polypeptides, labeling of one or the other Smad polypeptides, scintillation proximity, homogeneous time resolved fluorescence, fluorescence resonance energy transfer and fluorescence polarization.

- 22. A method of testing compounds, comprising the 10 steps of:
  - a) providing (i) a Smad polypeptide comprising the C-terminal domain, (ii) a polypeptide comprising the L45 loop of the kinase domain corresponding to a receptor of the TGF-β or BMP family, and (iii) a test compound; and
- b) contacting said Smad polypeptide with said receptor polypeptide under conditions where binding can take place, wherein said contacting is performed in the presence and absence of said compound; and
- c) detecting an increase or decrease in the binding of
   said Smad polypeptide to said kinase domain in the presence of said compound.
- 23. The method of claim 22, wherein the Smad polypeptide is selected from the group consisting of Smad1, Smad2,25 Smad3, Smad5 and Smad8.

24. The method of claim 22, wherein said drug may be used to treat ailments selected from the group consisting of pancreatic cancer, breast cancer, ovarian cancer, colon cancer, esophageal cancer, head and neck cancers, fibrosis of the kidney, fibrosis of the liver, fibrosis of the lung, Alzheimer's disease, memory loss, inflammation, wound healing, bone growth, immunoregulation, blood cell formation and atherosclerosis.

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25. The method of claim 22, wherein said detection is selected from the group consisting of solid support immobilization of one or the other polypeptides, labeling of one or the other polypeptides, scintillation proximity, homogeneous time resolved fluorescence, fluorescence resonance energy transfer and fluorescence polarization.

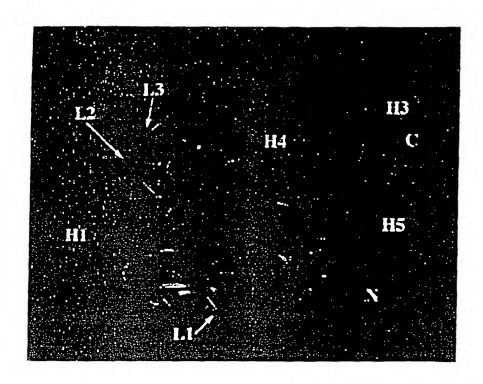


Figure 1

	552 (544 X D 10, 165 (544 X D 10, 16) (16) (16) (16) (16) (16) (16) (16)
397 397 397 397 397 397 403 403 403 397 503 503	552 465 424 424 465 418 819 839 876 876
O 0% - 60% buried  Productions  DIC-4  DIC-4  DIC-4  DIC-5  DIC-5  DIC-6  DIC-6  DIC-6  DIC-6  DIC-6  DIC-6  DIC-7	DPC.4 DAG 6 GREE STATE STATE STATE STATE DAGS GOODER COLORS STATE STATE STATE DAGS GOODER COLORS GOODER COLORS STATE STATE STATE DAGS SAAAGIGVDDLRR-CILCRASFVROWGEVRROWGESTRETPERTILLED CONTINUOUS TOWN TOWING SPINNDSS VS A VYOL TRMCTIRMSFVROWGE WORDOWTS TPCWIE LILL MGPLOWL TOWINGS PS - VRCS SMS A VYOL TRMCTIRMSFVROWGE WORDOWTS TPCWIE LILL MGPLOWL DWI TOWINGS PS - VRCS SWS A VYOL TRMCTIRMSFVROWGE WORDOWTS TPCWIE INLINGPLOWLD WITTOWIGS PS - VRCS SWS A VYOL TRMCTIRMSFVROWGE WORDOWTS TPCWIE INLINGPLOWLD WITTOWIGS PROPERSYS A VYOL TRMCTIRMSFVROWGE WORDOWTS TPCWIE INLINGPLOWLD WITTOWIGS PROPERSYS A VYOL TRMCTIRMSFVROWGE WORDOWTS TPCWIE INLINGPLOWLD RISS SWS A VYOL TRMCTIRMS FVROWGE WORDOWTS TPCWIE INLINGPLOWLD RISS SWS A VYOL TRMCTIRMS FVROWGE WORDOWTS TPCWIE LILL MAPLAYID WARROUT TOWING TPRINGPROWED THE NAPLAYID WARROUT TOWING TPCWIE LILL MAPLAYID WARROUT TOWING TAND TOWING TPCWIE LILL MAPLAYID WARROUT TOWING TWAND TAND TOWING THE LILL MAPLAYID WARROUT TOWING TWAND TAND TOWING THE LILL MAPLAYID WARROUT TOWING TWAND TAND TOWING THE LILL MAPLAYID WARROUT TOWING THE LILL MAPLAYING T
Conser Tumoria Davelog Smad4/UPC4 Smad1 Smad3 Smad3 Smad3 Smad3 Smad4/UPC4 Smad 2 Smad 2 Smad 3 Smad 2 Smad 3 Smad 4 Smad 3 Smad 4 Smad 3 Smad 3 Smad 3 Smad 3 Smad 3 Smad 3 Smad 4 Smad 3 Smad 4 Smad 5 Smad 4 Smad 5 Smad	Sixxid / OPCA Smad 1 Smad 2 Smad 3 Smad 3 Sma - 2 Sma - 3 Sma - 3

Figure 2

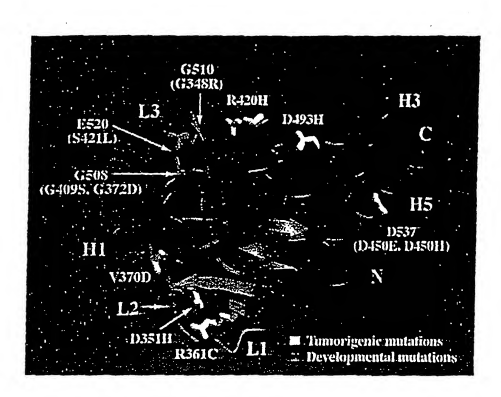
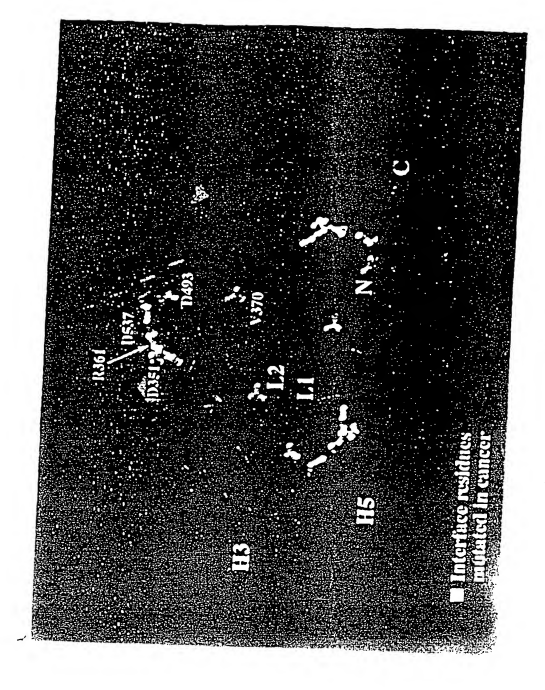


Figure 2b



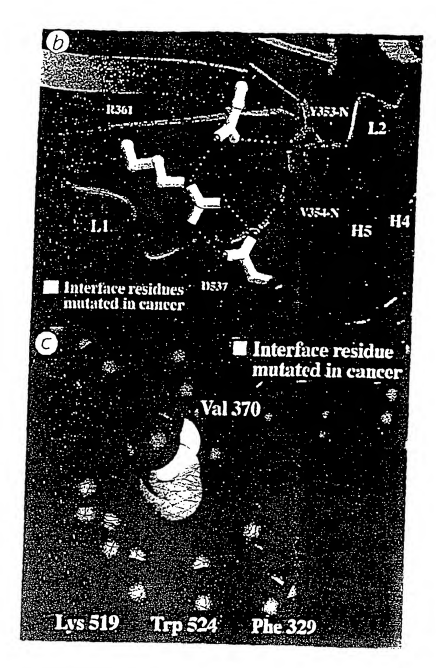


Figure 3 (cont.)

d.

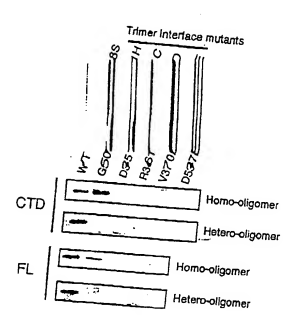
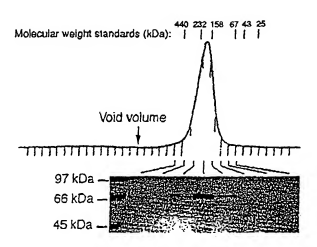


Figure 3d

а



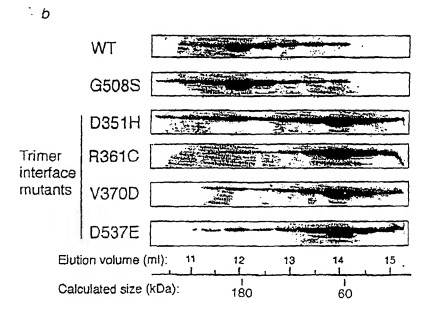
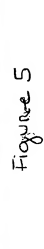
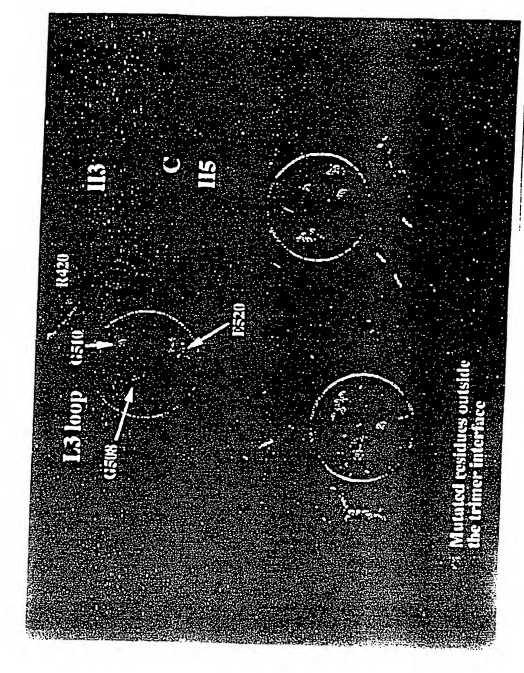
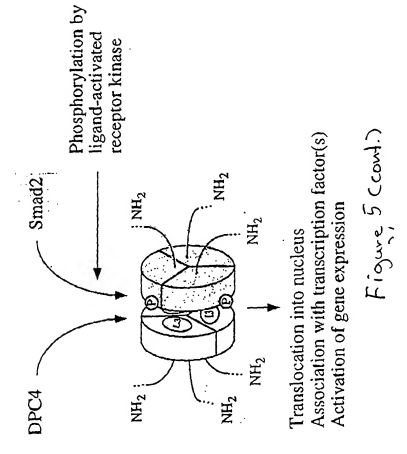


Figure 4



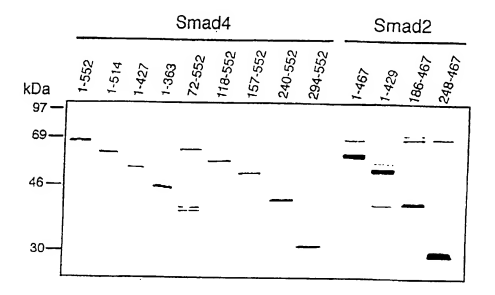




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a			
Smad4	Construct	Smad4 (1-552)	Smad2 (1 <b>-4</b> 67)
	1-552	+	+ ;
	1-499	+	-
	1-365	+	-
	1-148	-	-
	144-309	-	-
	310-552	+	+
Smad2	Construct	Smad2 (1-457)	Smad4 (1-552)
De la companya de la	1-467	+	+
	1-433	+	-
	1-319	-	-
	1-185	-	-
	187-269	-	-
	271-467	+	+

Figure 6



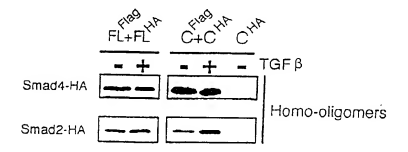
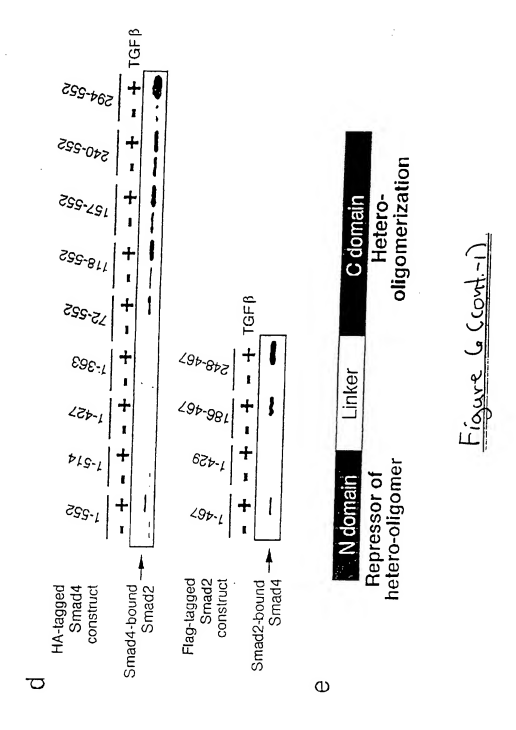


Figure 6 (cont.)



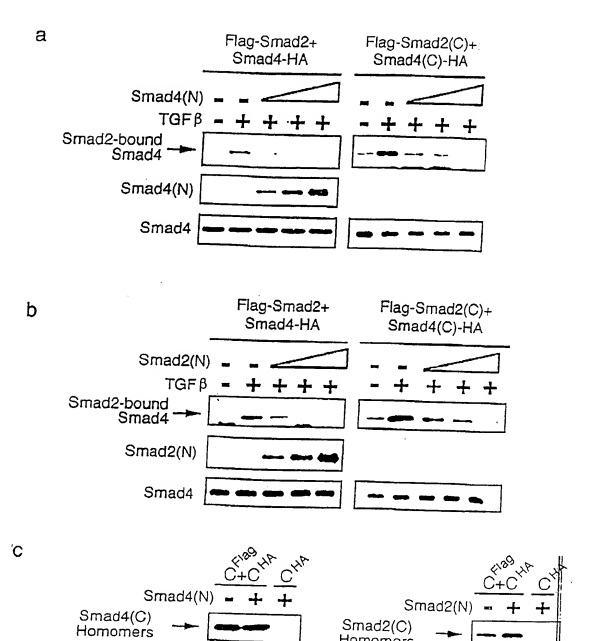


Figure 7

Smad2(C) Homomers

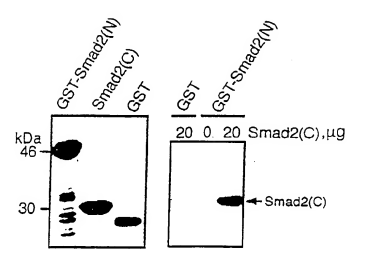


Figure 7 (cont)

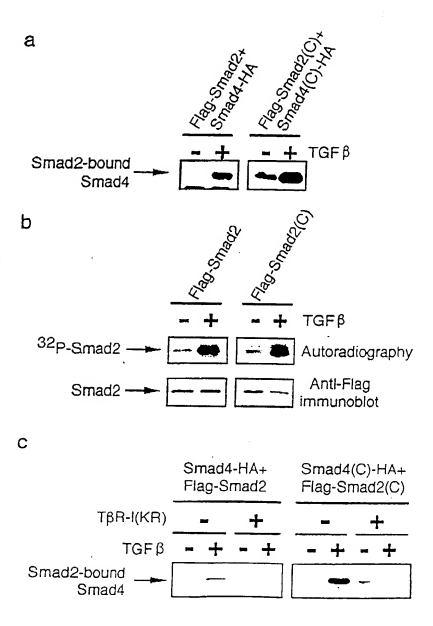
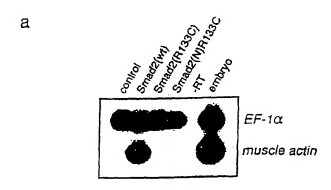
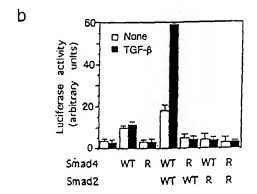


Figure 8

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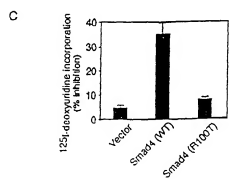


Figure 9



b

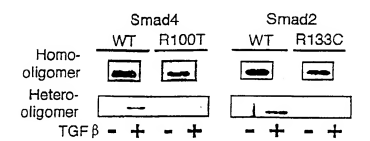


Figure 10

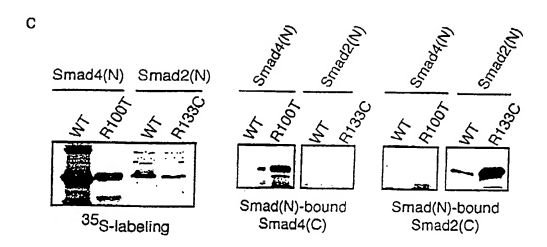
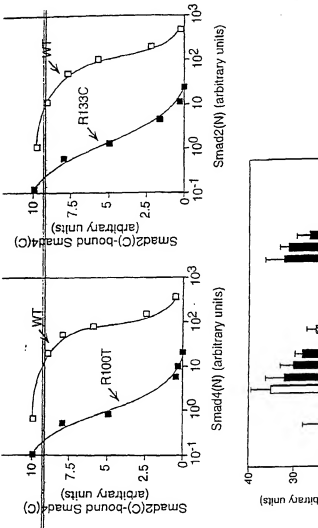
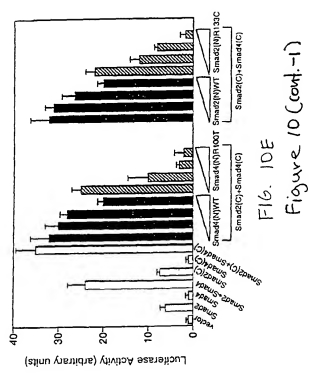


Figure 10 (cont.)





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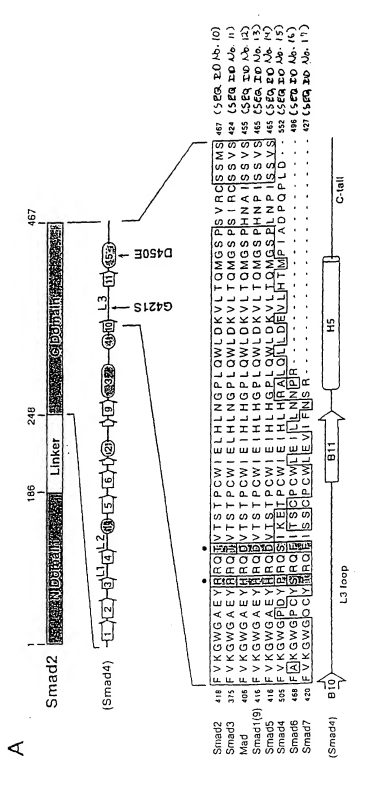


FIGURE 11A

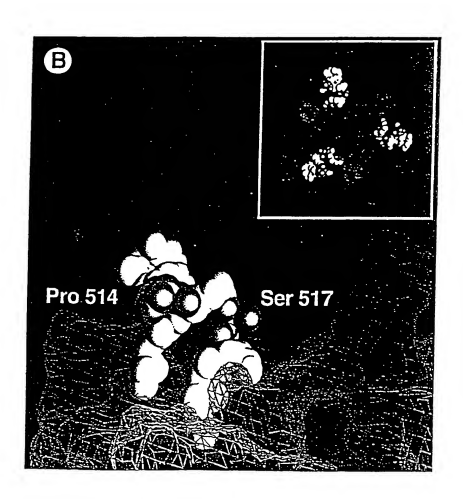


FIGURE 11B

A. Receptor Binding

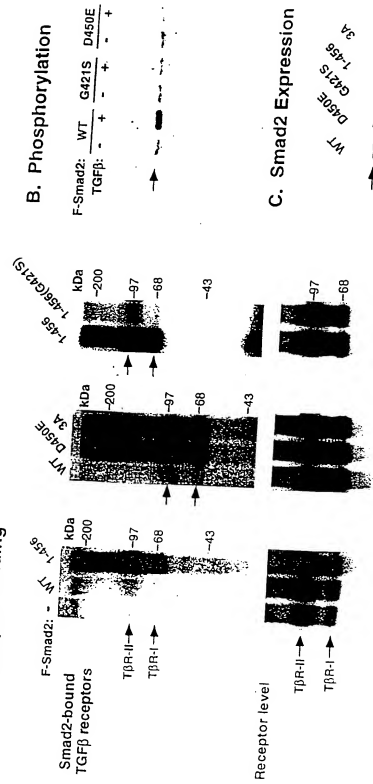


FIGURE 12

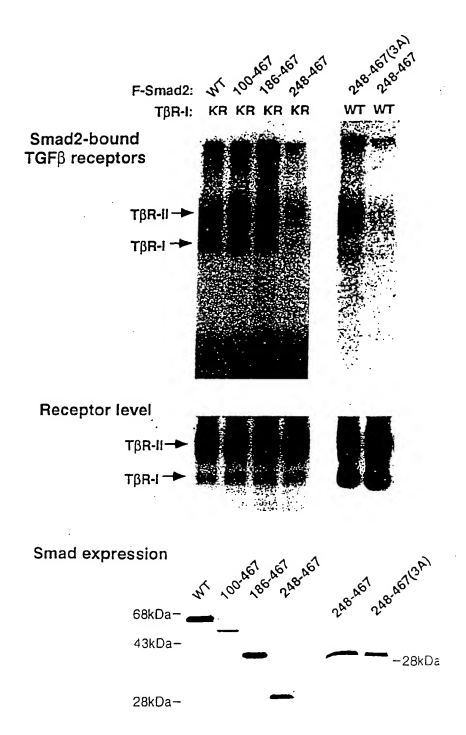


FIGURE 13

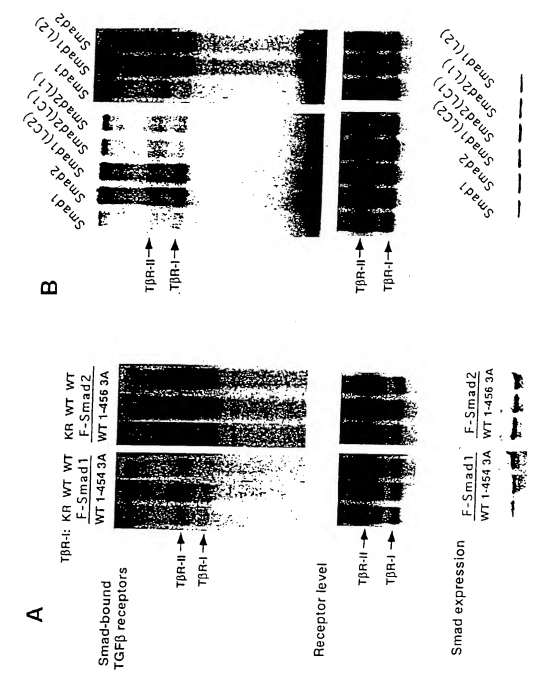
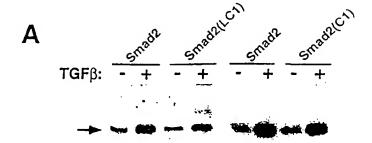
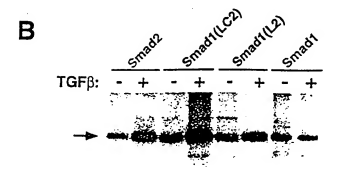
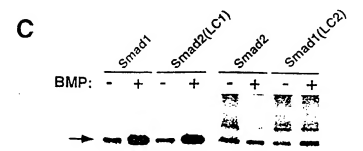


FIGURE 14







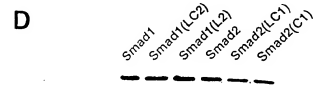


FIGURE 15

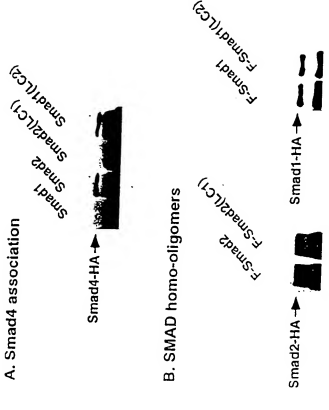


FIGURE 16

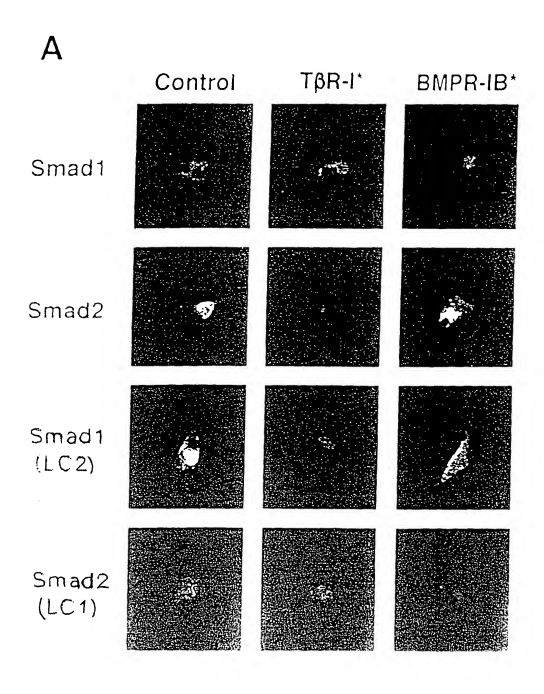
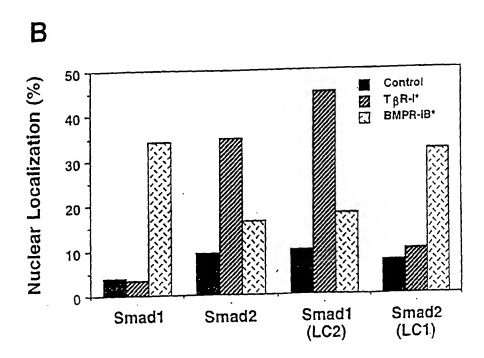
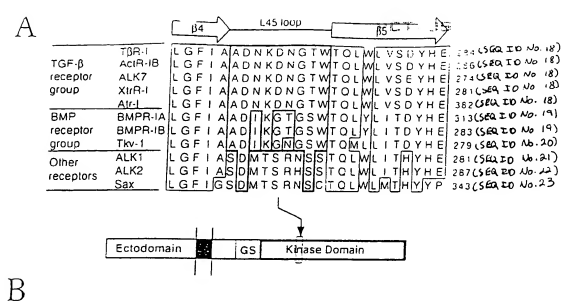


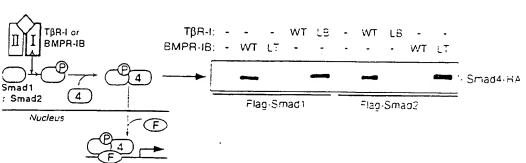
FIGURE 17A

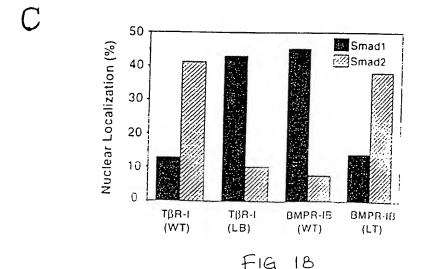
PCT/US98/13721

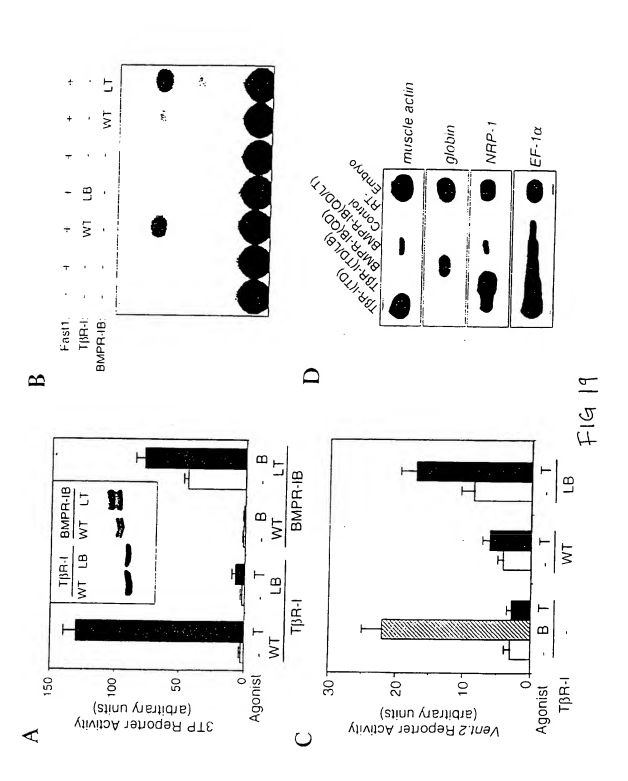


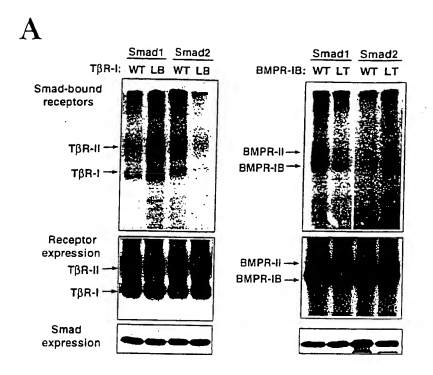
FRGURE 17B











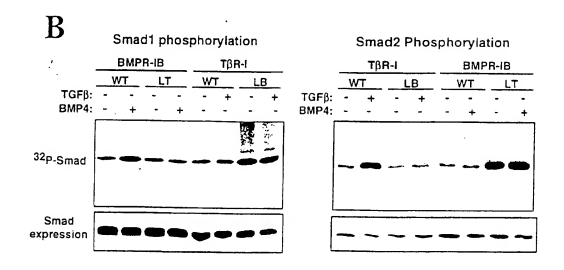
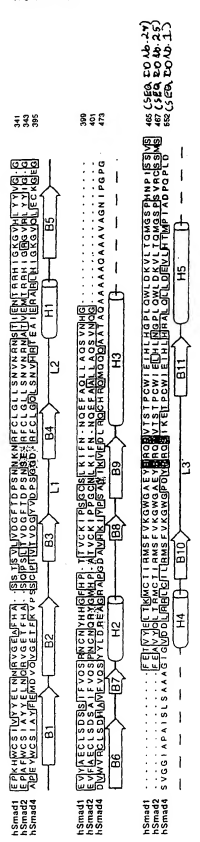
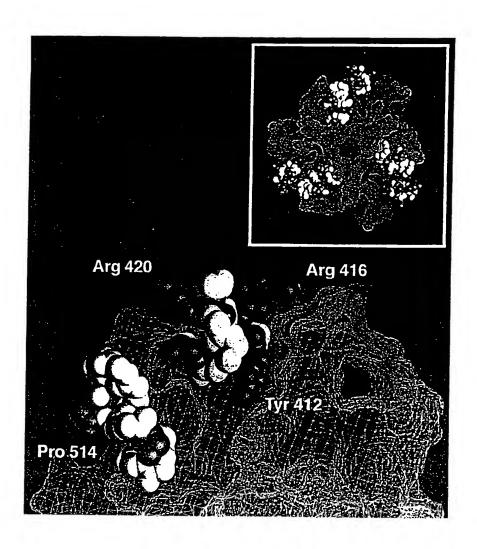


FIG 20

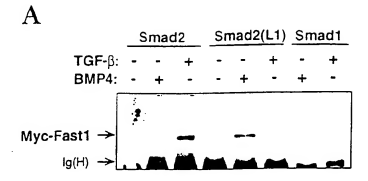




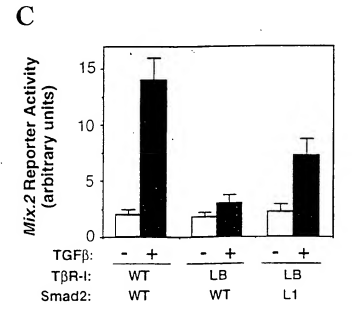
⋖



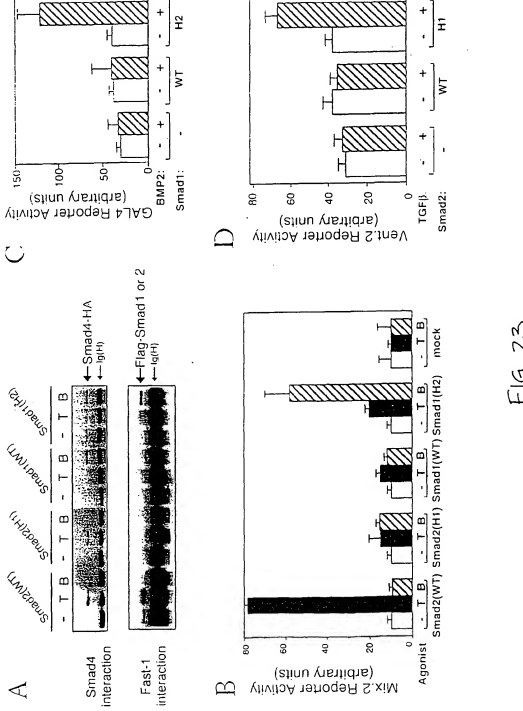
F19 21B

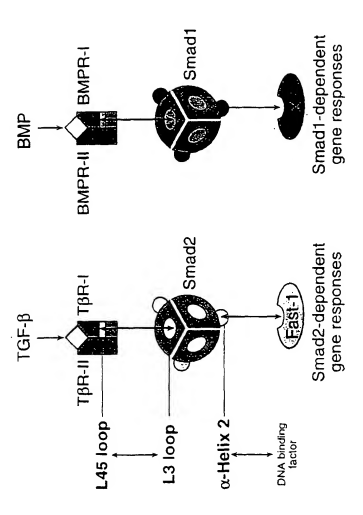






F16 22





F1G 24

International application No.
PCT/US98/13721

A. CLASSIFICATION OF SUBJECT MATTER  IPC(6) :G01N 33/53; C07K 14/51, 14/495, 14/71					
US CL	US CL :435/7.1; 530/399				
According to International Patent Classification (IPC) or to both national classification and IPC					
R. FIELDS SEARCHED					
Minimum d	ocumentation searched (classification system followed	by classification symbols)			
U.S. :	435/7.1				
Documentat	Documentation searched other than minimum documentation to the extent that such documents are included in the fields searched				
Electronic d	lats base consulted during the international search (na	me of data base and, where practicable	, search terms used)		
APS, CAPLUS search terms: smad, fastl, bmp, tgf-beta, MH2 domain, L3 loop, L45 loop, kinase domain					
C. DOC	UMENTS CONSIDERED TO BE RELEVANT				
Category*	Citation of document, with indication, where app	propriate, of the relevant passages	Relevant to claim No.		
Y	MASSAGUE et al. TGF-beta signallin Trends Cell Biol. May 1997, pages following: page 188, column 2; page 191; Figures 1-4.	187-192, see especially the	1-25		
Y	LIU et al. A human Mad protein acting as a BMP-regulated transcriptional activator. Nature. 13 June 1996, Vol. 381, pages 620-623, see especially the following: page 623, column 1, full paragraph 2.		18-21		
Y	CHEN et al. A transcriptional partner beta signalling. Nature. 24 October 696, see especially the following: Ab 694, paragraph bridging columns 1-2.	1996, Vol. 383, pages 691-	13-17		
X Further documents are listed in the continuation of Box C. See patent family annex.					
Special categories of cited documents:  "T"  later document published after the international filing date or priority date and not in conflict with the application but cited to understand the principle or theory underlying the invention					
*B* ex	to be of particular relevance  "B"  earlier document published on or after the international filing date  "X"  document of perticular relevance; the claimed invention cannot be considered novel or cannot be considered to involve an inventive step				
cit	scument which may throw doubts on priority claim(s) or which is ted to establish the publication date of another estation or other social reason (as specified)	"Y" document of particular relevance; th			
.0. qo	document referring to an oral disclosure, use, exhibition or other means combined with one or more other such documents, such combination being obvious to a person skilled in the art				
	pe document published prior to the international filing date but later than "a." document member of the same patent family the priority date claimed				
	Date of the actual completion of the international search  17 SEPTEMBER 1998  Date of mailing of the international search 22 OCT 1998				
Name and mailing address of the ISA/US Commissioner of Patents and Trademarks Box PCT Washington, D.C. 20231  Authorized officer DAVID S. ROMEO  FOR			/ R		
Facsimile 1	No. (703) 305-3230	Telephone No. (703) 308-0196			

International application No.
PCT/US98/13721

C (Continua	tion). DOCUMENTS CONSIDERED TO BE RELEVANT	
Category*	Citation of document, with indication, where appropriate, of the relevant passa	ges Relevant to claim No
Y	LAGNA et al. Partnership between DPC4 and SMAD proteir TGF-beta signalling pathways. Nature. 31 October 1996, Vo 383, pages 832-836, see especially the following: Figure 5; pa 836, column 1, full paragraph 1.	1.
Y	KRETZSCHMAR et al. The TGF-beta family mediator Smace phosphorylated directly and activated functionally by the BMI receptor kinase. Genes Dev., 15 April 1997, Vol. 11, No. 8, p. 984-995, see especially the following: page 992, paragraph bridging columns 1-2.	P
Y	ZHANG et al. Receptor-associated Mad homologues synergizeffectors of the TGF-beta response. Nature. 12 September 19 Vol. 383, pages 168-172, see sepecially the following: Abstrafigure 5.	96,
<b>4.</b>	FIELDS et al. A novel genetic system to detect protein-protein interactions. Nature. 20 July 1989, Vol. 340, pages 245-246.	n 1-25
<b>A</b> .	HELDIN C-H. Dimerization of cell surface receptors in signa transduction. Cell. 27 January 1995, Vol. 80, pages 213-223, page 219, column 2, last paragraph.	
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International application No. PCT/US98/13721

Box I Observations where certain claims were found unsearchable (Continuation of item 1 of first sheet)				
This international report has not been established in respect of certain claims under Article 17(2Xa) for the following reasons:				
1. Claims Nos.: because they relate to subject matter not required to be searched by this Authority, namely:				
Claims Nos.:  because they relate to parts of the international application that do not comply with the prescribed requirements to such an extent that no meaningful international search can be carried out, specifically:				
3. Claims Nos.; because they are dependent claims and are not drafted in accordance with the second and third sentences of Rule 6.4(a).				
Box II Observations where unity of invention is lacking (Continuation of item 2 of first sheet)				
This International Searching Authority found multiple inventions in this international application, as follows:				
Please See Extra Sheet.				
1. X As all required additional search fees were timely paid by the applicant, this international search report covers all searchable claims.				
2. As all searchable claims could be searched without effort justifying an additional fee, this Authority did not invite payment of any additional fee.				
3. As only some of the required additional search fees were timely paid by the applicant, this international search report covers only those claims for which fees were paid, specifically claims Nos.:				
4. No required additional search fees were timely paid by the applicant. Consequently, this international search report is restricted to the invention first mentioned in the claims; it is covered by claims Nos.:				
Remark on Protest  The additional search fees were accompanied by the applicant's protest.				
No protest accompanied the payment of additional search fees,				

International application No. PCT/US98/13721

BOX II. OBSERVATIONS WHERE UNITY OF INVENTION WAS LACKING This ISA found multiple inventions as follows:

This application contains the following inventions or groups of inventions which are not so linked as to form a single inventive concept under PCT Rule 13.1. In order for all inventions to be searched, the appropriate additional search fees must be paid.

Group I, claim(s) 1-4, drawn to a method of testing compounds wherein a Smad4 L3 loop and a complementary Smad polypeptide are employed.

Group II, claim(s) 5-8, drawn to a method of testing compounds wherein the C-terminal domain of two Smad polypeptides are employed.

Group III, claim(s) 9-12, drawn to a method of testing compounds wherein a C-terminal domain of a Smad polypeptide and the L45 loop of a kinase domain are employed.

Group IV, claim(s)13-17, drawn to a method of testing compounds wherein the alpha helix 2 of the SH2 domain of a Smad polypeptide and a DNA binding polypeptide are employed.

Group V, claim(s) 18-21, drawn to a method of testing compounds wherein two C-terminal domains from Smad polypeptides and a N-terminal domain of a Smad polypeptide are employed.

Group VI, claim(s) 22-25, drawn to a method of testing compounds wherein a Smad C-terminal domain and a L45 loop of a kinase domain.

The inventions listed as Groups I-VI do not relate to a single inventive concept under PCT Rule 13.1 because, under PCT Rule 13.2, they lack the same or corresponding special technical features for the following reasons:

With respect to unity of invention PCT Rule 13.1 states:

The international application shall relate to one invention only or to a group of inventions so linked as to form a single general inventive concept ("requirement of unity of invention").

### Additionally, PCT Rule 13.2 states:

Where a group of inventions is claimed in one and the same international application, the requirement of unity of invention referred to in Rule 13.1 shall be fulfilled only when there is a technical relationship among those inventions involving one or more of the same or corresponding special technical features. The expression "special technical features" shall mean those technical features that define a contribution which each of the claimed inventions, considered as a whole, makes over the prior art.

The claims recite a plurality of distinct methods involving the interaction Smad4 with other Smads, DNA binding proteins, or receptors, each method having different outcomes or process steps. The special technical feature of these methods is a Smad 4 polypeptide and its interaction with other proteins. In order for the methods to have unity of invention it is necessary that the special technical feature be a contribution over the prior art. However the prior art of MASSAGUE et al. discloses Smad 4 (Figure 2), Smads are directly phosphorylated by TGF-β family receptors and act by associating with other Smads and proteins that bind directly to the promoter regions of target genes (page 187, column 2). MASSAGUE et al. also suggest manipulation of the interactions of Smads with themselves, Smad4, TGF-β family receptors and DNA-binding components (page 191, paragraph bridging columns 1-2). Therefore the claimed inventions do not fulfill the requirements for unity of invention.

Accordingly, the claims are not so linked by a special technical feature within the meaning of PCT Rule 13.2 so as to form a single inventive concept.

With regard to the application of PCT Rule 13, 37 CFR § 1.475 concerning unity of invention states:

(d) If multiple products, processes of manufacture or uses are claimed, the first invention of the category first mentioned in the claims of the application and the first recited invention of each of the other categories related thereto will be considered as the main invention in the claims, see PCT Article 17(3)(a) and 37 C.F.R. § 1.476(c).

Pursuant to 37 C.F.R. § 1.475(d), this authority considers that where multiple products and processes are claimed, the first recited product, method of making a product, and method of using a product, together with the first recited of each of the other such inventions related thereto, shall constitute the main invention. Further, pursuant to 37 C.F.R. § 1.475(d) it considers that any subsequently recited products and/or methods constitute separate groups.

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	Accordingly, Groups I-VI constitute separate groups of inventions.
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